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Justin Whitle

#### BOX PATENT APPLICATION

**Assistant Commissioner for Patents** Washington, DC 20231

> U.S. Patent Application Entitled: METHODS AND COMPOSITIONS FOR THE RE: DIAGNOSIS AND TREATMENT OF CANCER - Gary L. Clayman (INGN:022)

Sir:

Transmitted herewith for filing is a 108-page patent specification including 25 claims and an abstract. Also included are Figures 1-7B on 8 sheets);. The specification and drawings constitute the application of Gary L. Clayman and Deborah Wilson for the captioned application.

The instant application claims priority to U.S. Serial No. 60/007,810 filed November 30, 1995 under 37 C.F.R. § 1.78(a)(3).

Also transmitted herewith is a diskette containing the computer-readable form of those sequences in the specification, a Statement as Required Under 37 C.F.R. § 1.821(f), and a separate paper copy of the sequence listing.

Assistant Commissioner for Patents November 27, 1996 Page 2

Please note that this application is filed without an inventors' Declaration and Assignment, a Declaration Claiming Small Entity Status, a Power of Attorney, and filing fees. Pursuant to 37 C.F.R. § 1.53(b) and (d), the Applicants request the Patent and Trademark Office to accept this application and accord a serial number and filing date as of the date this application is deposited with the U.S. Postal Service for Express Mail. Further, the Applicants request that the NOTICE OF MISSING PARTS-FILING DATE GRANTED pursuant to 37 C.F.R. § 1.53(d) be sent to the undersigned Applicants' representative.

Please date stamp and return the enclosed postcard to evidence receipt of this application.

Respectfully submitted,

Steven L. Highlander

Reg. No. 37,642

SLH/vp

Encl: Patent Application

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PATENT INGN:022



# METHODS AND COMPOSITIONS FOR THE DIAGNOSIS AND TREATMENT OF CANCER

By:

Gary L. Clayman

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Justin Whitley

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## **BACKGROUND OF THE INVENTION**

The present application is a continuation-in-part of U.S. Provisional Patent Application Serial No. 60/007,810 filed November 30, 1995. The entire text of the above-referenced disclosure is specifically incorporated by reference herein without disclaimer

#### A. Field of the Invention

The present invention is related generally to the field of cancer biology. In particular, the invention relates to compositions and methods of treatment for squamous cell carcinoma. Also provided is an animal model for the examination of microscopic residual tumors and tumor seeding into body cavities, as well as methods for treatment thereof.

#### B. Related Art

Balancing rates of cell proliferation and cell death is important in maintaining normal tissue homeostasis. Disruption of this balance may be a major factor in the multistep process of tumorigenesis, and inhibition of apoptosis, or programmed cell death, is one cause of this disruption. The effects of such defects are catastrophic, causing over half a million deaths annually in the United States alone.

There is considerable evidence implicating mutations of the *p53* gene, a tumor suppressor, in the etiology of many human cancers. Reports have demonstrated that growth of several different human cancer cell lines, including representatives of colon cancer, glioblastoma, breast cancer, osteosarcoma and lung cancer can be functionally suppressed by viral-mediated transfer of a wild-type *p53* gene. Induction of exogenous *p53* expression of wild-type *p53* has been shown to induce apoptosis in colon cancer cell lines and in human lung cancer spheroids, suggesting a role for *p53* in programmed cell death.

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Patients with squamous cell carcinoma of the head and neck (SCCHN) are afflicted with a disease that often has profound effects on speech, swallowing and cosmesis. Moreover, the overall rate of survival among these patients, approximately 50%, has remained unchanged for the nearly 30 years since contemporary surgery and radiation therapy were instituted. Recurrences are predominantly local and regional as opposed to systemic, indicating that microscopic residual carcinoma at the primary tumor site are the main cause of mortality. Given these facts, the ability to effectively attack microscopic residual disease in SCCHN is an endeavor that could improve the therapeutic efficacy of cancer treatments.

#### **SUMMARY OF THE INVENTION**

Therefore, it is an object of the present invention to provide improved methods for the *in vivo* treatment of squamous cell carcinoma. It is another object of the present invention to provide a method for assessing the development and treatment of microscopic residual carcinoma and microscopic tumor seeding of body cavities.

In fulfilling these objects, there is provided a method for treating a subject with squamous cell carcinoma comprising the steps of (a) providing an expression construct comprising a promoter functional in eukaryotic cells and a polynucleotide encoding a p53, wherein the polynucleotide is positioned sense to and under the control of the promoter; and (b) contacting the expression construct with the squamous cell carcinoma *in vivo*.

The squamous cell carcinoma may be a head and neck carcinoma. The endogenous p53 of the squamous cell carcinoma may or may not be mutated. The expression construct preferably is a viral vector, such as a retroviral vector, an adenoviral vector and an adeno-associated viral vector, with a replication-deficient adenoviral vector being most preferred. In a particular embodiment, the p53 gene is tagged so that expression of p53 from the expression vector can be detected. A preferred tag is an immunologic one, such as a continuous antibody epitope.

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The method may further comprise surgical resection of the tumor, with additional contacting of the tumor bed, or "artificial body cavity," with the expression construct after resection. The volume used to contact the tumor bed is about 3 ml. to about 10 ml. Where an adenovirus vector is employed, the amount of adenovirus administered in each contacting is about  $10^7$ ,  $10^8$ ,  $10^9$ ,  $10^{10}$ ,  $10^{11}$  or  $10^{12}$  pfu.

Continuous perfusion of the expression construct also is contemplated. The amount of construct delivered in continuous perfusion will be determined from the amount delivered via injections so as to approximate the same total dosage over a given time period, although somewhat greater total dosages may be achieved using continuous perfusion.

In another embodiment, the expression construct is injected into an natural body cavity such as the mouth, pharynx, esophagus, larynx, trachea, pleural cavity, peritoneal cavity, or hollow organ cavities including the bladder, colon or other visceral organs.

Also provided is a method for determining the effectiveness of a therapy on microscopic residual cancer comprising (a) providing a rodent with an incision into subcutaneous tissue; (b) seeding the incision with tumor cells; (c) treating the rodent with a therapeutic regimen; and (d) assessing the impact of the regimen on the development of tumors. The incision may be sealed following step (b) and prior to step (c). Further, the regimen may comprise introduction of a therapeutic composition into the incision, the incision being reopened after sealing and resealed after introduction of said therapeutic composition.

Other objects, features and advantages of the present invention will become apparent from the following detailed description. It should be understood, however, that the detailed description and the specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only, since various changes and modifications within the spirit

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and scope of the invention will become apparent to those skilled in the art from this detailed description.

#### **BRIEF DESCRIPTION OF THE DRAWINGS**

The following drawings form part of the present specification and are included to further demonstrate certain aspects of the present invention. The invention may be better understood by reference to one or more of these drawings in combination with the detailed description of specific embodiments presented herein:

- FIG. 1. Transduction efficiency of SCCHN cell lines Tu-138 (closed triangles) and Tu177 (closed squares). A recombinant  $\beta$ -gal adenovirus was used to infect the cells at different MOI ranging from 10 to 100. The percentages of  $\beta$ -gal-positive cells were obtained from scoring 500 cells each on replicate dishes.
- FIG. 2A and FIG. 2B. Inhibition of SCCHN cell growth *in vitro*. (FIG. 2A) Growth curve of mock-infected Tu-138 cells (closed circles), dl312-infected cells (closed triangles), and Ad5CMV-p53-infected cells (closed squares). (FIG. 2B) Growth curve of mock-infected Tu-177 cells (open circles), dl312-infected cells (open triangles), and Ad5CMV-p53-infected cells (open squares). At each indicated time point, three dishes of cells were trypsinized and counted. The mean  $\pm$  SEM of cell counts per triplicate wells following infection were plotted against the number of days since infection.
- FIG. 3A, FIG. 3B, FIG. 3C and FIG. 3D. Composite Growth Curve of four SCCHN cell lines. (FIG. 3A) Tu-138. (FIG. 3B) Tu-177. (FIG. 3C) MDA 686-LN. (FIG. 3D) MDA 886. Mock infected cells (closed circles), dl312-infected cells (closed triangle), and Ad5CMV-p53 infected cells (closed square). The mean of cell counts per triplicate wells following infection were plotted against the number of days since infection; bars, SEM.

- **FIG. 4.** Growth curve of normal fibroblast cell line. Mock-infected cells (closed circle), dl312-infected cells (closed triangle), and Ad5CMV-*p53*-infected cells (closed square).
- FIG. 5A and FIG. 5B. Composite growth curve of SCCHN cell lines. FIG. 5A: Tu-138; FIG. 5B: MDA 686LN; Mock infected cells (open square), dl312 infected cells (open triangle), and Ad5CMV-p53 infected cells (open circle). At each indicated time point, three dishes of cells were trypsinized and counted. The mean of cell counts per triplicate dishes were plotted against the number of hours post-infection; bars, SEM.

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FIG. 6A and FIG. 6B. Labeling of DNA breaks in apoptotic cells with biotinylated-dUTP by TUNEL method. Following infection, flow cytometric analysis for apoptosis was performed in a time course study. FIG. 6A. Tu-138 cells which are infected with dl312, a replication-defective adenovirus (panel 1-panel 4), Tu-138 cells infected with the wild-type p53 adenovirus (panel 5- panel 8). FIG. 6B. MDA 686LN cells which are infected with dl312, a replication-defective adenovirus (A-D), MDA 686LN cells infected with the wild-type p53 adenovirus (E-H). Ap stands for apoptosis.

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FIG. 7A and FIG. 7B. Composite growth curve of SCCHN cell line. FIG. 7A: Tu138; FIG. 7B: MD686LN; Mock infected cells (open circles), dl312 infected cells (closed triangles), Ad5CMV-p53 infected cells (open squares) and Ad5CMV-p53-FLAG infected cells (closed squares). At each indicated time point, three dishes of cells were trypsinized and counted. The mean of cell counts per triplicate dishes were plotted against the number of hours post-infection; bars, SEM.

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#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Information available to date suggests that one of the primary shortcomings of treatments for SCCHN is the inability to completely eradicate disease at the primary tumor site, or in the immediate local or regional tissues. Therefore, the present invention is designed to provide gene therapeutic methodologies that permit a more complete and effective treatment of SCCHN, specifically, by attacking microscopic residual carcinoma. This methodology may be used alone or as an adjunct to more conventional treatments such as chemo- or radiotherapy or surgical intervention. Moreover, using an animal model specifically designed to address microscopic residual carcinoma, as well as microscopic tumor seeding of body cavities, the present inventor has demonstrated efficacy of these methods. The details of the invention are described more completely below.

More generally, it now has been observed that p53 gene therapy of cancers may be effective regardless of the p53 status of the tumor cell. Surprisingly, therapeutic effects have been observed when a viral vector carrying the wild-type p53 gene is used to treat a tumor, the cells of which express a functional p53 molecule. This result would not have been predicted based on the current understanding of how tumor suppressors function. It also is surprising given that normal cells, which also express a functional p53 molecule, are apparently unaffected by expression of high levels of p53 from a viral construct. This raises the possibility that p53 gene therapy may be more broadly applicable to the treatment of cancers than was initially suspected.

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#### A. p53 Proteins and Polynucleotides

Throughout this application, the term "p53" is intended to refer to the exemplified p53 molecules as well as all p53 homologues from other species. "Wild-type" and "mutant" p53 refer, respectively, to a p53 gene expressing normal tumor suppressor activity and to a p53 gene lacking or having reduced suppressor activity and/or having transforming activity. Thus "mutant" p53's are not merely sequence variants but rather, are those variants showing altered functional profiles.

p53 is currently recognized as a tumor suppressor gene (Montenarh, 1992). High levels have been found in many cells transformed by chemical carcinogenesis, ultraviolet radiation, and several viruses, including SV40. The p53 gene is a frequent target of mutational inactivation in a wide variety of human tumors and is already documented to be the most frequently-mutated gene in common human cancers (Mercer, 1992). It is mutated in over 50% of human NSCLC (Hollestein et al., 1991) and in a wide spectrum of other tumors.

While tumors containing a mutated *p53* gene are a preferred target according to the present invention, the utility of the claimed *p53* expression vectors extends to the treatment of tumors having wild-type or functional *p53*. Though the mechanism is not completely understood, the present inventor has determined that *p53* expression can limit the growth of tumors expressing a functional *p53* product, and even induce apoptosis in such cells. Thus, the *p53* status of a tumor, though potentially useful in a diagnostic context, is not essential for practice of the present invention. This phenomenon is not limited to SCCHN tumors, but may be applied to a wide variety of malignancies including gliomas, sarcomas, carcinomas, leukemias, lymphomas and melanoma, including tumors of the skin, liver, testes, bone, brain, pancreas, head and neck, stomach, liver, lung, ovary, breast, colon, prostate and bladder.

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#### p53 Polypeptides

The p53 gene encodes a 375-amino-acid phosphoprotein that can form complexes with viral proteins such as large-T antigen and E1B. The protein is found in normal tissues and cells, but at concentrations which are minute by comparison with many transformed cells or tumor tissue. Interestingly, wild-type p53 appears to be important in regulating cell growth and division. Overexpression of wild-type p53 has been shown in some cases to be anti-proliferative in human tumor cell lines. Thus p53 can act as a negative regulator of cell growth (Weinberg, 1991) and may directly suppress uncontrolled cell growth or indirectly activate genes that suppress this growth. Thus, absence or inactivation of wild-type p53 may contribute to transformation. However, some studies indicate that the presence of mutant p53 may be necessary for full expression of the transforming potential of the gene.

Although wild-type p53 is recognized as a centrally important growth regulator in many cell types, its genetic and biochemical traits appear to have a role as well. Mis-sense mutations are common for the p53 gene and are essential for the transforming ability of the oncogene. A single genetic change prompted by a point mutation can create carcinogenic p53. Unlike other oncogenes, however, p53 point mutations are known to occur in at least 30 distinct codons, often creating dominant alleles that produce shifts in cell phenotype without a reduction to homozygosity. Additionally, many of these dominant negative alleles appear to be tolerated in the organism and passed on in the germ line. Various mutant alleles appear to range from minimally dysfunctional to strongly penetrant, dominant negative alleles (Weinberg, 1991).

Casey and colleagues have reported that transfection of DNA encoding wild-type p53 into two human breast cancer cell lines restores growth suppression control in such cells (Casey et al., 1991). A similar effect has also been demonstrated on transfection of wild-type, but not mutant, p53 into human lung cancer cell lines (Takahasi et al., 1992). The p53 wild-type appears dominant over the mutant gene and will select against proliferation when transfected into cells with the mutant gene. Expression of the transfected p53 does not affect the growth of

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normal cells with endogenous p53. Thus, such constructs might be taken up by normal cells without adverse effects.

It is thus possible that the treatment of p53-associated cancers with wild-type p53 may reduce the number of malignant cells. However, studies such as those described above are far from achieving such a goal, not least because DNA transfection cannot be employed to introduce DNA into cancer cells within a patients' body.

#### p53-Encoding Polynucleotides

The polynucleotides according to the present invention may encode an entire *p53* gene, a functional *p53* protein domain, or any *p53* polypeptide. "Complementary" polynucleotides are those which are capable of base-pairing according to the standard Watson-Crick complementarity rules. That is, the larger purines will base pair with the smaller pyrimidines to form combinations of guanine paired with cytosine (G:C) and adenine paired with either thymine (A:T) in the case of DNA, or adenine paired with uracil (A:U) in the case of RNA. Inclusion of less common bases such as inosine, 5-methylcytosine, 6-methyladenine, hypoxanthine and others in hybridizing sequences does not interfere with pairing.

As used herein, the term "complementary sequences" means polynucleotide sequences that are substantially complementary over their entire length and have very few base mismatches. For example, sequences of fifteen bases in length may be termed complementary when they have a complementary nucleotide at thirteen or fourteen positions. Naturally, sequences which are "completely complementary" will be sequences which are entirely complementary throughout their entire length and have no base mismatches.

Other sequences with lower degrees of homology also are contemplated. For example, an antisense construct which has limited regions of high homology, but also contains a non-homologous region (e.g., a ribozyme) could be designed. These molecules, though having less than 50% homology, would bind to target sequences under appropriate conditions.

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The polynucleotides may be derived from genomic DNA, *i.e.*, cloned directly from the genome of a particular organism. In other embodiments, however, the polynucleotides may be complementary DNA (cDNA). cDNA is DNA prepared using messenger RNA (mRNA) as template. Thus, a cDNA does not contain any interrupted coding sequences and usually contains almost exclusively the coding region(s) for the corresponding protein. In other embodiments, the polynucleotide may be produced synthetically.

It may be advantageous to combine portions of the genomic DNA with cDNA or synthetic sequences to generate specific constructs. For example, where an intron is desired in the ultimate construct, a genomic clone will need to be used. Introns may be derived from other genes in addition to p53. The cDNA or a synthesized polynucleotide may provide more convenient restriction sites for the remaining portion of the construct and, therefore, would be used for the rest of the sequence.

The human and mouse DNA sequence for p53 is provided in SEQ ID NO:1 and SEQ ID NO:3 respectively, with the corresponding amino acids being provided in SEQ ID NO:2 and SEQ ID NO:4, respectively.

It is contemplated that natural variants of p53 exist that have different sequences than those disclosed herein. Thus, the present invention is not limited to use of the provided polynucleotide sequence for p53 but, rather, includes use of any naturally-occurring variants. The present invention also encompasses chemically synthesized mutants of these sequences.

Another kind of sequence variant results from codon variation. Because there are several codons for most of the 20 normal amino acids, many different DNA's can encode the p53. Reference to the following table will allow such variants to be identified.

TABLE 1

Amino Acids			Codons					
Alanine	Ala	A	GCA	GCC	GCG	GCU		
Cysteine	Cys	C	UGC	UGU				
Aspartic acid	Asp	D	GAC	GAU				
Glutamic acid	Glu	E	GAA	GAG				
Phenylalanine	Phe	F	UUC	UUU				
Glycine	Gly	G	GGA	GGC	GGG	GGU		
Histidine	His	H	CAC	CAU				
Isoleucine	Ile	I	AUA	AUC	AUU			
Lysine	Lys	K	AAA	AAG				
Leucine	Leu	L	UUA	UUG	CUA	CUC	CUG	CUU
Methionine	Met	M	AUG					
Asparagine	Asn	N	AAC	AAU				
Proline	Pro	P	CCA	CCC	CCG	CCU		
Glutamine	Gln	Q	CAA	CAG				
Arginine	Arg	R	AGA	AGG	CGA	CGC	CGG	CGU
Serine	Ser	S	AGC	AGU	UCA	UCC	UCG	UCU
Threonine	Thr	T	ACA	ACC	ACG	ACU		
Valine	Val	V	GUA	GUC	GUG	GUU		
Tryptophan	Trp	W	UGG					
Tyrosine	Tyr	Y	UAC U.	AU				ir.

Allowing for the degeneracy of the genetic code, sequences that have between about 50% and about 75%, or between about 76% and about 99% of nucleotides that are identical to the nucleotides disclosed herein will be preferred. Sequences that are within the scope of "a p53-encoding polynucleotide" are those that are capable of base-pairing with a polynucleotide segment set forth above under intracellular conditions.

As stated above, although the *p53* encoding sequences may be full length genomic or cDNA copies, or large fragments thereof. The present invention also may employ shorter oligonucleotides of p53. Sequences of 17 bases long should occur only once in the human genome and, therefore, suffice to specify a unique target sequence. Although shorter oligomers are easier to make and increase *in vivo* accessibility, numerous other factors are involved in determining the specificity of base-pairing. Both binding affinity and sequence specificity of an oligonucleotide to its complementary target increases with increasing length. It is contemplated that oligonucleotides of 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20 base pairs will be used, for example, in the preparation of p53 mutants and in PCR reactions.

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Any sequence of 17 bases long should occur only once in the human genome and, therefore, suffice to specify a unique target sequence. Although shorter oligomers are easier to make and increase *in vivo* accessibility, numerous other factors are involved in determining the specificity of hybridization. Both binding affinity and sequence specificity of an oligonucleotide to its complementary target increases with increasing length.

In certain embodiments, one may wish to employ constructs which include other elements, for example, those which include C-5 propyne pyrimidines. Oligonucleotides which contain C-5 propyne analogues of uridine and cytidine have been shown to bind RNA with high affinity (Wagner *et al.*, 1993).

It also is well understood by the skilled artisan that, inherent in the definition of a biologically functional equivalent protein or peptide, is the concept that there is a limit to the number of changes that may be made within a defined portion of the molecule and still result in a molecule with an acceptable level of equivalent biological activity. Biologically functional equivalent peptides are thus defined herein as those peptides in which certain, not most or all, of the amino acids may be substituted. In particular, where the N-terminus of the p16 protein is concerned, it is contemplated that only about 16 or more preferably, about 5 amino acids may be

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changed within a given peptide. Of course, a plurality of distinct proteins/peptides with different substitutions may easily be made and used in accordance with the invention.

Amino acid substitutions are generally based on the relative similarity of the amino acid side-chain substituents, for example, their hydrophobicity, hydrophilicity, charge, size, and the like. An analysis of the size, shape and type of the amino acid side-chain substituents reveals that arginine, lysine and histidine are all positively charged residues; that alanine, glycine and serine are all a similar size; and that phenylalanine, tryptophan and tyrosine all have a generally similar shape. Therefore, based upon these considerations, arginine, lysine and histidine; alanine, glycine and serine; and phenylalanine, tryptophan and tyrosine; are defined herein as biologically functional equivalents.

In making changes, the hydropathic index of amino acids may be considered. Each amino acid has been assigned a hydropathic index on the basis of their hydrophobicity and charge characteristics, these are: isoleucine (+4.5); valine (+4.2); leucine (+3.8); phenylalanine (+2.8); cysteine/cystine (+2.5); methionine (+1.9); alanine (+1.8); glycine (-0.4); threonine (-0.7); serine (-0.8); tryptophan (-0.9); tyrosine (-1.3); proline (-1.6); histidine (-3.2); glutamate (-3.5); glutamine (-3.5); aspartate (-3.5); asparagine (-3.5); lysine (-3.9); and arginine (-4.5).

The importance of the hydropathic amino acid index in conferring interactive biological function on a protein is generally understood in the art (Kyte & Doolittle, 1982, incorporated herein by reference). It is known that ceratin amino acids may be substituted for other amino acids having a similar hydropathic index or score and still retain a similar biological activity. In making changes based upon the hydropathic index, the substitution of amino acids whose hydropathic indices are within  $\pm 2$  is preferred, those which are within  $\pm 1$  are particularly preferred, and those within  $\pm 0.5$  are even more particularly preferred.

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It is understood that an amino acid can be substituted for another having a similar hydrophilicity value and still obtain a biologically equivalent protein. As detailed in U.S. Patent 4,554,101, the following hydrophilicity values have been assigned to amino acid residues: arginine (+3.0); lysine (+3.0); aspartate (+3.0  $\pm$  1); glutamate (+3.0  $\pm$  1); serine (+0.3); asparagine (+0.2); glutamine (+0.2); glycine (0); threonine (-0.4); proline (-0.5  $\pm$  1); alanine (-0.5); histidine (-0.5); cysteine (-1.0); methionine (-1.3); valine (-1.5); leucine (-1.8); isoleucine (-1.8); tyrosine (-2.3); phenylalanine (-2.5); tryptophan (-3.4).

In making changes based upon similar hydrophilicity values, the substitution of amino acids whose hydrophilicity values are within  $\pm 2$  is preferred, those which are within  $\pm 1$  are particularly preferred, and those within  $\pm 0.5$  are even more particularly preferred.

#### **B.** Expression Vectors

Throughout this application, the term "expression construct" is meant to include any type of genetic construct containing a nucleic acid coding for a gene product in which part or all of the nucleic acid encoding sequence is capable of being transcribed. The transcript may be translated into a protein, but it need not be. Thus, in certain embodiments, expression includes both transcription of a p53 gene and translation of a p53 mRNA into a p53 protein product. In other embodiments, expression only includes transcription of the nucleic acid encoding a p53 or its complement.

In order for the construct to effect expression of at least a *p53* transcript, the polynucleotide encoding the *p53* polynucleotide will be under the transcriptional control of a promoter. A "promoter" refers to a DNA sequence recognized by the synthetic machinery of the host cell, or introduced synthetic machinery, that is required to initiate the specific transcription of a gene. The phrase "under transcriptional control" means that the promoter is in the correct location in relation to the polynucleotide to control RNA polymerase initiation and expression of the polynucleotide.

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to fix the place of initiation.

The term promoter will be used here to refer to a group of transcriptional control modules that are clustered around the initiation site for RNA polymerase II. Much of the thinking about how promoters are organized derives from analyses of several viral promoters, including those for the HSV thymidine kinase (tk) and SV40 early transcription units. These studies, augmented by more recent work, have shown that promoters are composed of discrete functional modules, each consisting of approximately 7-20 bp of DNA, and containing one or more recognition sites for transcriptional activator or repressor proteins.

At least one module in each promoter functions to position the start site for RNA synthesis. The best known example of this is the TATA box, but in some promoters lacking a TATA box, such as the promoter for the mammalian terminal deoxynucleotidyl transferase gene and the promoter for the SV40 late genes, a discrete element overlying the start site itself helps

Additional promoter elements regulate the frequency of transcriptional initiation. Typically, these are located in the region 30-110 bp upstream of the start site, although a number of promoters have recently been shown to contain functional elements downstream of the start site as well. The spacing between promoter elements frequently is flexible, so that promoter function is preserved when elements are inverted or moved relative to one another. In the tk promoter, the spacing between promoter elements can be increased to 50 bp apart before activity begins to decline. Depending on the promoter, it appears that individual elements can function either co-operatively or independently to activate transcription.

The particular promoter that is employed to control the expression of a p53 polynucleotide is not believed to be critical, so long as it is capable of expressing the polynucleotide in the targeted cell at sufficient levels. Thus, where a human cell is targeted, it is preferable to position the polynucleotide coding region adjacent to and under the control of a

promoter that is capable of being expressed in a human cell. Generally speaking, such a promoter might include either a human or viral promoter.

In various embodiments, the human cytomegalovirus (CMV) immediate early gene promoter, the SV40 early promoter and the Rous sarcoma virus long terminal repeat can be used to obtain high-level expression of the *p53* polynucleotide. The use of other viral or mammalian cellular or bacterial phage promoters which are well-known in the art to achieve expression of polynucleotides is contemplated as well, provided that the levels of expression are sufficient to produce a growth inhibitory effect.

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By employing a promoter with well-known properties, the level and pattern of expression of a polynucleotide following transfection can be optimized. For example, selection of a promoter which is active in specific cells, such as tyrosinase (melanoma), alpha-fetoprotein and albumin (liver tumors), CC10 (lung tumor) and prostate-specific antigen (prostate tumor) will permit tissue-specific expression of p53 polynucleotides. Table 2 lists several elements/promoters which may be employed, in the context of the present invention, to regulate the expression of p53 constructs. This list is not intended to be exhaustive of all the possible elements involved in the promotion of p53 expression but, merely, to be exemplary thereof.

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Enhancers were originally detected as genetic elements that increased transcription from a promoter located at a distant position on the same molecule of DNA. This ability to act over a large distance had little precedent in classic studies of prokaryotic transcriptional regulation. Subsequent work showed that regions of DNA with enhancer activity are organized much like promoters. That is, they are composed of many individual elements, each of which binds to one or more transcriptional proteins.

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The basic distinction between enhancers and promoters is operational. An enhancer region as a whole must be able to stimulate transcription at a distance; this need not be true of a promoter region or its component elements. On the other hand, a promoter must have one or

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more elements that direct initiation of RNA synthesis at a particular site and in a particular orientation, whereas enhancers lack these specificities. Promoters and enhancers are often overlapping and contiguous, often seeming to have a very similar modular organization.

Additionally any promoter/enhancer combination (as per the Eukaryotic Promoter Data Base EPDB) could also be used to drive expression of a *p53* construct. Use of a T3, T7 or SP6 cytoplasmic expression system is another possible embodiment. Eukaryotic cells can support cytoplasmic transcription from certain bacteriophage promoters if the appropriate bacteriophage polymerase is provided, either as part of the delivery complex or as an additional genetic expression vector.

# TABLE 2

ENHANCER	
Immunoglobulin Heavy Chain	
Immunoglobulin Light Chain	
T-Cell Receptor	
HLA DQ $\alpha$ and DQ $\beta$	
ß-Interferon	
Interleukin-2	
Interleukin-2 Receptor	
MHC Class II 5	
MHC Class II HLA-DRα	
ß-Actin	
Muscle Creatine Kinase	
Prealbumin (Transthyretin)	
Elastase I	
Metallothionein	
Collagenase	
Albumin Gene	
α-Fetoprotein	
τ-Globin	
B-Globin	
c-fos	
c-HA-ras	
Insulin	
Neural Cell Adhesion Molecule (NCAM)	
$\alpha_1$ -Antitrypsin	

ENHANCER			
H2B (TH2B) Histone			
Mouse or Type I Collagen			
Glucose-Regulated Proteins (GRP94 and GRP78)			
Rat Growth Hormone			
Human Serum Amyloid A (SAA)			
Troponin I (TN I)			
Platelet-Derived Growth Factor			
Duchenne Muscular Dystrophy			
SV40			
Polyoma			
Retroviruses			
Papilloma Virus			
Hepatitis B Virus			
Human Immunodeficiency Virus			
Cytomegalovirus			
Gibbon Ape Leukemia Virus			

Further, selection of a promoter that is regulated in response to specific physiologic signals can permit inducible expression of the p53 construct. For example, with the polynucleotide under the control of the human PAI-1 promoter, expression is inducible by tumor necrosis factor.

5 Table 3 illustrates several promoter/inducer combinations:

TABLE 3

Element	Inducer
MT II	Phorbol Ester (TFA) Heavy metals
MMTV (mouse mammary tumor virus)	Glucocorticoids
B-Interferon	poly(rI)X poly(rc)
Adenovirus 5 <u>E2</u>	Ela
c-jun	Phorbol Ester (TPA), H <sub>2</sub> O <sub>2</sub>
Collagenase	Phorbol Ester (TPA)
Stromelysin	Phorbol Ester (TPA), IL-1
SV40	Phorbol Ester (TPA)
Murine MX Gene	Interferon, Newcastle Disease Virus
GRP78 Gene	A23187
α-2-Macroglobulin	IL-6
Vimentin	Serum
MHC Class I Gene H-2kB	Interferon
HSP70	Ela, SV40 Large T Antigen
Proliferin	Phorbol Ester-TPA
Tumor Necrosis Factor	FMA
Thyroid Stimulating Hormone $\alpha$ Gene	Thyroid Hormone

In certain embodiments of the invention, the delivery of an expression vector in a cell may be identified *in vitro* or *in vivo* by including a marker in the expression vector. The marker would result in an identifiable change to the transfected cell permitting easy identification of expression. Usually the inclusion of a drug selection marker aids in cloning and in the selection of transformants. Alternatively, enzymes such as herpes simplex virus thymidine kinase (tk) (eukaryotic) or chloramphenicol acetyltransferase (CAT) (prokaryotic) may be employed.

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Immunologic markers also can be employed. The selectable marker employed is not believed to be important, so long as it is capable of being expressed along with the polynucleotide encoding p53. Further examples of selectable markers are well known to one of skill in the art.

One typically will include a polyadenylation signal to effect proper polyadenylation of the transcript. The nature of the polyadenylation signal is not believed to be crucial to the successful practice of the invention, and any such sequence may be employed. The inventor has employed the SV40 polyadenylation signal in that it was convenient and known to function well in the target cells employed. Also contemplated as an element of the expression construct is a terminator. These elements can serve to enhance message levels and to minimize read through from the construct into other sequences.

In preferred embodiments of the present invention, the expression construct comprises a virus or engineered construct derived from a viral genome. The ability of certain viruses to enter cells via receptor-mediated endocytosis and, in some cases, integrate into the host cell chromosomes, have made them attractive candidates for gene transfer in to mammalian cells. However, because it has been demonstrated that direct uptake of naked DNA, as well as receptor-mediated uptake of DNA complexes (discussed below), expression vectors need not be viral but, instead, may be any plasmid, cosmid or phage construct that is capable of supporting expression of encoded genes in mammalian cells, such as pUC or Bluescript<sup>TM</sup> plasmid series.

#### **Retroviruses**

The retroviruses are a group of single-stranded RNA viruses characterized by an ability to convert their RNA to double-stranded DNA in infected cells by a process of reverse-transcription (Coffin, 1990). The resulting DNA then stably integrates into cellular chromosomes as a provirus and directs synthesis of viral proteins. The integration results in the retention of the viral gene sequences in the recipient cell and its descendants. The retroviral genome contains three genes - gag, pol, and env - that code for capsid proteins, polymerase enzyme, and envelope components, respectively. A sequence found upstream from the gag

gene, termed  $\Psi$ , functions as a signal for packaging of the genome into virions. Two long terminal repeat (LTR) sequences are present at the 5' and 3' ends of the viral genome. These contain strong promoter and enhancer sequences and are also required for integration in the host cell genome (Coffin, 1990).

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In order to construct a retroviral vector, a nucleic acid encoding a p53 is inserted into the viral genome in the place of certain viral sequences to produce a virus that is replication-defective. In order to produce virions, a packaging cell line containing the gag, pol and env genes but without the LTR and vomponents is constructed (Mann et al., 1983). When a recombinant plasmid containing a human cDNA, together with the retroviral LTR and vequences is introduced into this cell line (by calcium phosphate precipitation for example), the vequence allows the RNA transcript of the recombinant plasmid to be packaged into viral particles, which are then secreted into the culture media (Nicolas and Rubenstein, 1988; Temin, 1986; Mann et al., 1983). The media containing the recombinant retroviruses is then collected, optionally concentrated, and used for gene transfer. Retroviral vectors are able to infect a broad variety of cell types. However, integration and stable expression require the division of host cells (Paskind et al., 1975).

A novel approach designed to allow specific targeting of retrovirus vectors was recently developed based on the chemical modification of a retrovirus by the chemical addition of lactose residues to the viral envelope. This modification could permit the specific infection of hepatocytes via sialoglycoprotein receptors.

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A different approach to targeting of recombinant retroviruses was designed in which biotinylated antibodies against a retroviral envelope protein and against a specific cell receptor were used. The antibodies were coupled via the biotin components by using streptavidin (Roux et al., 1989). Using antibodies against major histocompatibility complex class I and class II antigens, they demonstrated the infection of a variety of human cells that bore those surface antigens with an ecotropic virus in vitro (Roux et al., 1989).

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#### Adenoviruses

Human adenoviruses are double-stranded DNA tumor viruses with genome sizes of approximate 36 kb (Tooze, 1981). As a model system for eukaryotic gene expression, adenoviruses have been widely studied and well characterized, which makes them an attractive system for development of adenovirus as a gene transfer system. This group of viruses is easy to grow and manipulate, and they exhibit a broad host range *in vitro* and *in vivo*. In lytically infected cells, adenoviruses are capable of shutting off host protein synthesis, directing cellular machineries to synthesize large quantities of viral proteins, and producing copious amounts of virus.

The E1 region of the genome includes E1A and E1B which encode proteins responsible for transcription regulation of the viral genome, as well as a few cellular genes. E2 expression, including E2A and E2B, allows synthesis of viral replicative functions, *e.g.* DNA-binding protein, DNA polymerase, and a terminal protein that primes replication. E3 gene products prevent cytolysis by cytotoxic T cells and tumor necrosis factor and appear to be important for viral propagation. Functions associated with the E4 proteins include DNA replication, late gene expression, and host cell shutoff. The late gene products include most of the virion capsid proteins, and these are expressed only after most of the processing of a single primary transcript from the major late promoter has occurred. The major late promoter (MLP) exhibits high efficiency during the late phase of the infection (Stratford-Perricaudet and Perricaudet, 1991a).

As only a small portion of the viral genome appears to be required *in cis* (Tooze, 1981), adenovirus-derived vectors offer excellent potential for the substitution of large DNA fragments when used in connection with cell lines such as 293 cells. Ad5-transformed human embryonic kidney cell lines (Graham, *et al.*, 1977) have been developed to provide the essential viral proteins *in trans*. The inventor thus reasoned that the characteristics of adenoviruses rendered them good candidates for use in targeting cancer cells *in vivo* (Grunhaus & Horwitz, 1992).

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Particular advantages of an adenovirus system for delivering foreign proteins to a cell include (i) the ability to substitute relatively large pieces of viral DNA by foreign DNA; (ii) the structural stability of recombinant adenoviruses; (iii) the safety of adenoviral administration to humans; and (iv) lack of any known association of adenoviral infection with cancer or malignancies; (v) the ability to obtain high titers of the recombinant virus; and (vi) the high infectivity of Adenovirus.

Further advantages of adenovirus vectors over retroviruses include the higher levels of gene expression. Additionally, adenovirus replication is independent of host gene replication, unlike retroviral sequences. Because adenovirus transforming genes in the E1 region can be readily deleted and still provide efficient expression vectors, oncogenic risk from adenovirus vectors is thought to be negligible (Grunhaus & Horwitz, 1992).

In general, adenovirus gene transfer systems are based upon recombinant, engineered adenovirus which is rendered replication-incompetent by deletion of a portion of its genome, such as E1, and yet still retains its competency for infection. Sequences encoding relatively large foreign proteins can be expressed when additional deletions are made in the adenovirus genome. For example, adenoviruses deleted in both E1 and E3 regions are capable of carrying up to 10 Kb of foreign DNA and can be grown to high titers in 293 cells (Stratford-Perricaudet and Perricaudet, 1991a). Surprisingly persistent expression of transgenes following adenoviral infection has also been reported.

Adenovirus-mediated gene transfer has recently been investigated as a means of mediating gene transfer into eukaryotic cells and into whole animals. For example, in treating mice with the rare recessive genetic disorder ornithine transcarbamylase (OTC) deficiency, it was found that adenoviral constructs could be employed to supply the normal OTC enzyme. Unfortunately, the expression of normal levels of OTC was only achieved in 4 out of 17 instances (Stratford-Perricaudet *et al.*, 1991b). Therefore, the defect was only partially corrected

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in most of the mice and led to no physiological or phenotypic change. These type of results therefore offer little encouragement for the use of adenoviral vectors in cancer therapy.

Attempts to use adenovirus to transfer the gene for cystic fibrosis transmembrane conductance regulator (CFTR) into the pulmonary epithelium of cotton rats have also been partially successful, although it has not been possible to assess the biological activity of the transferred gene in the epithelium of the animals (Rosenfeld *et al.*, 1992). Again, these studies demonstrated gene transfer and expression of the CFTR protein in lung airway cells but showed no physiologic effect. In the 1991 *Science* article, Rosenfeld *et al.* showed lung expression of a1-antitrypsin protein but again showed no physiologic effect. In fact, they estimated that the levels of expression that they observed were only about 2% of the level required for protection of the lung in humans, *i.e.*, far below that necessary for a physiologic effect.

The gene for human  $\alpha$ 1-antitrypsin has been introduced into the liver of normal rats by intraportal injection, where it was expressed and resulted in the secretion of the introduced human protein into the plasma of these rats (Jaffe *et al.*, 1992). However, and disappointingly, the levels that were obtained were not high enough to be of therapeutic value.

These type of results do not demonstrate that adenovirus is able to direct the expression of sufficient protein in recombinant cells to achieve a physiologically relevant effect, and they do not, therefore, suggest a usefulness of the adenovirus system for use in connection with cancer therapy. Furthermore, prior to the present invention, it was thought that p53 could not be incorporated into a packaging cell, such as those used to prepare adenovirus, as it would be toxic. As E1B of adenovirus binds to p53, this was thought to be a further reason why adenovirus and p53 technology could not be combined.

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## Other Viral Vectors as Expression Constructs

Other viral vectors may be employed as expression constructs in the present invention. Vectors derived from viruses such as vaccinia virus (Ridgeway, 1988; Baichwal and Sugden, 1986; Coupar *et al.*, 1988) adeno-associated virus (AAV) (Ridgeway, 1988; Baichwal and Sugden, 1986; Hermonat and Muzycska, 1984) and herpesviruses may be employed. They offer several attractive features for various mammalian cells (Friedmann, 1989; Ridgeway, 1988; Baichwal and Sugden, 1986; Coupar *et al.*, 1988; Horwich *et al.*, 1990).

With the recent recognition of defective hepatitis B viruses, new insight was gained into the structure-function relationship of different viral sequences. *in vitro* studies showed that the virus could retain the ability for helper-dependent packaging and reverse transcription despite the deletion of up to 80% of its genome (Horwich *et al.*, 1990). This suggested that large portions of the genome could be replaced with foreign genetic material. The hepatotropism and persistence (integration) were particularly attractive properties for liver-directed gene transfer. Chang *et al.* recently introduced the chloramphenical acetyltransferase (CAT) gene into duck hepatitis B virus genome in the place of the polymerase, surface, and pre-surface coding sequences. It was cotransfected with wild-type virus into an avian hepatoma cell line. Culture media containing high titers of the recombinant virus were used to infect primary duckling hepatocytes. Stable CAT gene expression was detected for at least 24 days after transfection (Chang *et al.*, 1991).

# C. Alternative Methods for Gene Delivery

In order to effect expression of *p53* constructs, the expression vector must be delivered into a cell. As described above, the preferred mechanism for delivery is via viral infection where the expression vector is encapsidated in an infectious adenovirus particle.

Several non-viral methods for the transfer of expression vectors into cultured mammalian cells also are contemplated by the present invention. These include calcium phosphate

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precipitation (Graham and Van Der Eb, 1973; Chen and Okayama, 1987; Rippe et al., 1990) DEAE-dextran (Gopal, 1985), electroporation (Tur-Kaspa et al., 1986; Potter et al., 1984), direct microinjection (Harland and Weintraub, 1985), DNA-loaded liposomes (Nicolau and Sene, 1982; Fraley et al., 1979) and lipofectamine-DNA complexes, cell sonication (Fechheimer et al., 1987), gene bombardment using high velocity microprojectiles (Yang et al., 1990), polycations (Boussif et al., 1995) and receptor-mediated transfection (Wu and Wu, 1987; Wu and Wu, 1988). Some of these techniques may be successfully adapted for in vivo or ex vivo use.

In one embodiment of the invention, the adenoviral expression vector may simply consist of naked recombinant vector. Transfer of the construct may be performed by any of the methods mentioned above which physically or chemically permeabilize the cell membrane. For example, Dubensky *et al.* (1984) successfully injected polyomavirus DNA in the form of CaPO<sub>4</sub> precipitates into liver and spleen of adult and newborn mice demonstrating active viral replication and acute infection. Benvenisty and Neshif (1986) also demonstrated that direct intraperitoneal injection of CaPO<sub>4</sub> precipitated plasmids results in expression of the transfected genes. It is envisioned that DNA encoding an *p53* construct may also be transferred in a similar manner *in vivo*.

Another embodiment of the invention for transferring a naked DNA expression vector into cells may involve particle bombardment. This method depends on the ability to accelerate DNA coated microprojectiles to a high velocity allowing them to pierce cell membranes and enter cells without killing them (Klein et al., 1987). Several devices for accelerating small particles have been developed. One such device relies on a high voltage discharge to generate an electrical current, which in turn provides the motive force (Yang et al., 1990). The microprojectiles used have consisted of biologically inert substances such as tungsten or gold beads.

Selected organs including the liver, skin, and muscle tissue of rats and mice have been bombarded in vivo (Yang et al., 1990; Zelenin et al., 1991). This may require surgical exposure

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of the tissue or cells, to eliminate any intervening tissue between the gun and the target organ. DNA encoding a p53 construct may be delivered via this method.

In a further embodiment of the invention, the expression vector may be entrapped in a liposome. Liposomes are vesicular structures characterized by a phospholipid bilayer membrane and an inner aqueous medium. Multilamellar liposomes have multiple lipid layers separated by aqueous medium. They form spontaneously when phospholipids are suspended in an excess of aqueous solution. The lipid components undergo self-rearrangement before the formation of closed structures and entrap water and dissolved solutes between the lipid bilayers (Ghosh and Bachhawat, 1991). Also contemplated are lipofectamine-DNA complexes.

Liposome-mediated polynucleotide delivery and expression of foreign DNA *in vitro* has been very successful. Wong *et al.* (1980) demonstrated the feasibility of liposome-mediated delivery and expression of foreign DNA in cultured chick embryo, HeLa and hepatoma cells. Nicolau *et al.* (1987) accomplished successful liposome-mediated gene transfer in rats after intravenous injection.

In certain embodiments of the invention, the liposome may be complexed with a hemagglutinating virus (HVJ). This has been shown to facilitate fusion with the cell membrane and promote cell entry of liposome-encapsulated DNA (Kaneda *et al.*, 1989). In other embodiments, the liposome may be complexed or employed in conjunction with nuclear non-histone chromosomal proteins (HMG-1) (Kato *et al.*, 1991). In yet further embodiments, the liposome may be complexed or employed in conjunction with both HVJ and HMG-1. In that such expression vectors have been successfully employed in transfer and expression of a polynucleotide *in vitro* and *in vivo*, then they are applicable for the present invention. Where a bacteriophage promoter is employed in the DNA construct, it also will be desirable to include within the liposome an appropriate bacteriophage polymerase.

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Another mechanism for transferring expression vectors into cells is receptor-mediated delivery. This approach takes advantage of the selective uptake of macromolecules by receptor-mediated endocytosis in almost all eukaryotic cells. Because of the cell type-specific distribution of various receptors, the delivery can be highly specific (Wu and Wu, 1993). Receptor-mediated gene targeting vehicles generally consist of two components: a cell receptor-specific ligand and a DNA-binding agent. Several ligands have been used for receptor-mediated gene transfer. The most extensively characterized ligands are asialoorosomucoid (ASOR) (Wu and Wu, 1987) and transferrin (Wagner *et al.*, 1993). Recently, a synthetic neoglycoprotein, which recognizes the same receptor as ASOR, has been used as a gene delivery vehicle (Ferkol *et al.*, 1993; Perales *et al.*, 1994) and epidermal growth factor (EGF) has also been used to deliver genes to squamous carcinoma cells (Myers, EPO 0273085).

In other embodiments, the delivery vehicle may comprise a ligand and a liposome. For a galactose-terminal employed lactosyl-ceramide, (1987)Nicolau et al. example, asialganglioside, incorporated into liposomes and observed an increase in the uptake of the insulin gene by hepatocytes. Thus, it is feasible that an adenoviral expression vector also may be specifically delivered into a cell type such as lung, epithelial or tumor cells, by any number of receptor-ligand systems, with or without liposomes. For example, epidermal growth factor (EGF) may be used as the receptor for mediated delivery of p53 construct in many tumor cells that exhibit upregulation of EGF receptor. Mannose can be used to target the mannose receptor on liver cells. Also, antibodies to CD5 (CLL), CD22 (lymphoma), CD25 (T-cell leukemia) and MAA (melanoma) can similarly be used as targeting moieties.

In certain embodiments, gene transfer may more easily be performed under *ex vivo* conditions. *Ex vivo* gene therapy refers to the isolation of cells from an animal, the delivery of a polynucleotide into the cells, *in vitro*, and then the return of the modified cells back into an animal. This may involve the surgical removal of tissue/organs from an animal or the primary culture of cells and tissues. Anderson *et al.*, U.S. Patent 5,399,346, and incorporated herein in

its entirety, disclose *ex vivo* therapeutic methods. During *ex vivo* culture, the expression vector can express the *p53* construct. Finally, the cells may be reintroduced into the original animal, or administered into a distinct animal, in a pharmaceutically acceptable form by any of the means described below.

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# D. Pharmaceutical Compositions and Routes of Administration

Where clinical application of an adenoviral expression vector according to the present invention is contemplated, it will be necessary to prepare the complex as a pharmaceutical composition appropriate for the intended application. Generally this will entail preparing a pharmaceutical composition that is essentially free of pyrogens, as well as any other impurities that could be harmful to humans or animals. One also will generally desire to employ appropriate salts and buffers to render the complex stable and allow for complex uptake by target cells.

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Aqueous compositions of the present invention comprise an effective amount of the expression vector, dissolved or dispersed in a pharmaceutically acceptable carrier or aqueous medium. Such compositions also are referred to as inocula. The phrases "pharmaceutically or pharmacologically acceptable" refer to molecular entities and compositions that do not produce an adverse, allergic or other untoward reaction when administered to an animal, or a human, as appropriate. As used herein, "pharmaceutically acceptable carrier" includes any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents and the like. The use of such media and agents for pharmaceutical active substances is well known in the art. Except insofar as any conventional media or agent is incompatible with the active ingredient, its use in the therapeutic compositions is contemplated. Supplementary active ingredients also can be incorporated into the compositions.

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Solutions of the active compounds as free base or pharmacologically acceptable salts can be prepared in water suitably mixed with a surfactant, such as hydroxypropylcellulose.

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Dispersions also can be prepared in glycerol, liquid polyethylene glycols, mixtures thereof and in oils. Under ordinary conditions of storage and use, these preparations contain a preservative to prevent the growth of microorganisms.

The expression vectors and delivery vehicles of the present invention may include classic pharmaceutical preparations. Administration of therapeutic compositions according to the present invention will be via any common route so long as the target tissue is available via that route. This includes oral, nasal, buccal, rectal, vaginal or topical. Alternatively, administration will be by orthotopic, intradermal, intraocular, subcutaneous, intramuscular, intraperitoneal or intravenous injection. Such compositions would normally be administered as pharmaceutically acceptable compositions that include physiologically acceptable carriers, buffers or other excipients.

The therapeutic compositions of the present invention are advantageously administered in the form of injectable compositions either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid prior to injection may also be prepared. These preparations also may be emulsified. A typical composition for such purpose comprises a pharmaceutically acceptable carrier. For instance, the composition may contain 10 mg, 25 mg, 50 mg or up to about 100 mg of human serum albumin per milliliter of phosphate buffered Other pharmaceutically acceptable carriers include aqueous solutions, non-toxic saline. excipients, including salts, preservatives, buffers and the like. Examples of non-aqueous solvents are propylene glycol, polyethylene glycol, vegetable oil and injectable organic esters Aqueous carriers include water, alcoholic/aqueous solutions, saline such as ethyloleate. solutions, parenteral vehicles such as sodium chloride, Ringer's dextrose, etc. Intravenous vehicles include fluid and nutrient replenishers. Preservatives include antimicrobial agents, anti-oxidants, chelating agents and inert gases. The pH and exact concentration of the various components the pharmaceutical composition are adjusted according to well known parameters.

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Additional formulations are suitable for oral administration. Oral formulations include such typical excipients as, for example, pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate and the like. The compositions take the form of solutions, suspensions, tablets, pills, capsules, sustained release formulations or powders. When the route is topical, the form may be a cream, ointment, salve, liquid or spray.

An effective amount of the therapeutic agent is determined based on the intended goal, for example (i) inhibition of tumor cell proliferation or (ii) elimination of tumor cells. The term "unit dose" refers to physically discrete units suitable for use in a subject, each unit containing a predetermined-quantity of the therapeutic composition calculated to produce the desired responses, discussed above, in association with its administration, i.e., the appropriate route and treatment regimen. The quantity to be administered, both according to number of treatments and unit dose, depends on the subject to be treated, the state of the subject and the protection desired. Precise amounts of the therapeutic composition also depend on the judgment of the practitioner and are peculiar to each individual.

In certain embodiments, it may be desirable to provide a continuous supply of therapeutic compositions to the patient. For intravenous or intraarterial routes, this is accomplished by drip system. For topical applications, repeated application would be employed. For various approaches, delayed release formulations could be used that provided limited but constant amounts of the therapeutic agent over and extended period of time. For internal application, continuous perfusion of the region of interest may be preferred. This could be accomplished by catheterization, post-operatively in some cases, followed by continuous administration of the therapeutic agent. The time period for perfusion would be selected by the clinician for the particular patient and situation, but times could range from about 1-2 hours, to 2-6 hours, to about 6-10 hours, to about 10-24 hours, to about 1-2 days, to about 1-2 weeks or longer. Generally, the dose of the therapeutic composition via continuous perfusion will be equivalent to that given by single or multiple injections, adjusted for the period of time over which the

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injections are administered. It is believed that higher doses may be achieved via perfusion, however.

#### **Clinical Protocol for SCCHN**

A clinical protocol has been developed to facilitate the treatment of SCCHN disease using the adenoviral constructs discussed below in the examples. In accordance with this protocol, patients having histologic proof of squamous cell carcinoma of the head and neck will be selected. Patients may, but need not have received previous chemo-, radio- or gene therapies. Optimally, patients will have adequate bone marrow function (defined as peripheral absolute granulocyte count of  $> 2,000/\text{mm}^3$  and platelet count of  $100,000/\text{mm}^3$ ), adequate liver function (bilirubin  $\leq 1.5 \text{ mg/dl}$ ) and adequate renal function (creatinine < 1.5 mg/dl).

The protocol calls for single dose administration, via intratumoral injection, of a pharmaceutical composition containing between  $10^6$  and  $10^9$  infectious particles of a p53 adenovirus expression construct. For tumors of  $\geq 4$  cm, the volume administered will be 4-10 ml (preferably 10 ml), while for tumors < 4 cm, a volume of 1-3 ml will be used (preferably 3 ml). Multiple injections will be delivered for a single dose, in 0.1-0.5 ml volumes, with spacing of approximately 1 cm or more.

The treatment course consists of about six doses, delivered over two weeks. Upon election by the clinician, the regimen may be continued, six doses each two weeks, or on a less frequent (monthly, bimonthly, quarterly, etc.) basis.

Where patients are eligible for surgical resection, the tumor will be treated as described above for at least two consecutive two-week treatment courses. Within one week of completion of the second (or more, e.g., third, fourth, fifth, sixth, seventh, eighth, etc.) course, the patient will receive surgical resection. Prior to close of the incision, 10 ml of a pharmaceutical composition containing the p53 adenovirus expression construct ( $10^6$ - $10^9$  infectious particles) will be delivered to the surgical site (operative bed) and allowed to remain in contact for at least 60

minutes. The wound is closed and a drain or catheter placed therein. On the third postoperative day, additional 10 ml of the pharmaceutical composition is administered via the drain and allowed to remain in contact with the operative bed for at least two hours. Removal by suction is then performed, and the drain removed at a clinically appropriate time.

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#### **Treatment of Artificial and Natural Body Cavities**

One of the prime sources of recurrent SCCHN is the residual, microscopic disease that remains at the primary tumor site, as well as locally and regionally, following tumor excision. In addition, there are analogous situations where natural body cavities are seeded by microscopic tumor cells. The effective treatment of such microscopic disease would present a significant advance in therapeutic regimens.

Thus, in certain embodiments, a cancer may be removed by surgical excision, creating a "cavity." Both at the time of surgery, and thereafter (periodically or continuously), the therapeutic composition of the present invention is administered to the body cavity. This is, in essence, a "topical" treatment of the surface of the cavity. The volume of the composition should be sufficient to ensure that the entire surface of the cavity is contacted by the expression construct.

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In one embodiment, administration simply will entail injection of the therapeutic composition into the cavity formed by the tumor excision. In another embodiment, mechanical application via a sponge, swab or other device may be desired. Either of these approaches can be used subsequent to the tumor removal as well as during the initial surgery. In still another embodiment, a catheter is inserted into the cavity prior to closure of the surgical entry site. The cavity may then be continuously perfused for a desired period of time.

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In another form of this treatment, the "topical" application of the therapeutic composition is targeted at a natural body cavity such as the mouth, pharynx, esophagus, larynx, trachea, pleural cavity, peritoneal cavity, or hollow organ cavities including the bladder, colon or other

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visceral organs. In this situation, there may or may not be a significant, primary tumor in the cavity. The treatment targets microscopic disease in the cavity, but incidentally may also affect a primary tumor mass if it has not been previously removed or a pre-neoplastic lesion which may be present within this cavity. Again, a variety of methods may be employed to affect the "topical" application into these visceral organs or cavity surfaces. For example, the oral cavity in the pharynx may be affected by simply oral swishing and gargling with solutions. However, topical treatment within the larynx and trachea may require endoscopic visualization and topical delivery of the therapeutic composition. Visceral organs such as the bladder or colonic mucosa may require indwelling catheters with infusion or again direct visualization with a cystoscope or other endoscopic instrument. Cavities such as the pleural and peritoneal cavities may be accessed by indwelling catheters or surgical approaches which provide access to those areas.

# Monitoring p53 Expression Following Administration

Another aspect of the present invention involves the monitoring of p53 expression following administration of the therapeutic composition. Because destruction of microscopic tumor cells cannot be observed, it is important to determine whether the target site has been effectively contacted with the expression construct. This may be accomplished by identifying cells in which the expression construct is actively producing the p53 product. It is important, however, to be able to distinguish between the exogenous p53 and that present in tumor and non-tumor cells in the treatment area. Tagging of the exogenous p53 with a tracer element would provide definitive evidence for expression of that molecule and not an endogenous version thereof.

One such tracer is provided by the FLAG biosystem (Hopp et al., 1988). The FLAG polypeptide is an octapeptide (AspTyrLysAspAspAspAspLys) and its small size does not disrupt the expression of the delivered gene therapy protein. The coexpression of FLAG and the protein of interest is traced through the use of antibodies raised against FLAG protein.

Other immunologic marker systems, such as the 6XHis system (Qiagen) also may be employed. For that matter, any linear epitope could be used to generate a fusion protein with p53 so long as (i) the immunologic integrity of the epitope is not compromised by the fusion and (ii) the functional integrity of p53 is not compromised by the fusion.

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# E. Combined Therapy Protocols

Tumor cell resistance to DNA damaging agents represents a major problem in clinical oncology. One goal of current cancer research is to find ways to improve the efficacy of chemo-and radiotherapy by combining it with gene therapy. For example, the herpes simplex-thymidine kinase (HS-tK) gene, when delivered to brain tumors by a retroviral vector system, successfully induced susceptibility to the antiviral agent ganciclovir (Culver, *et al.*, 1992). In the context of the present invention, it is contemplated that *p53* therapy could be used similarly in conjunction with chemo- or radiotherapeutic intervention.

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To kill cells, such as malignant or metastatic cells, using the methods and compositions of the present invention, one would generally contact a "target" cell with an expression vector and at least one DNA damaging agent. These compositions would be provided in a combined amount effective to kill or inhibit proliferation of the cell. This process may involve contacting the cells with the expression vector and the DNA damaging agent(s) or factor(s) at the same time. This may be achieved by contacting the cell with a single composition or pharmacological formulation that includes both agents, or by contacting the cell with two distinct compositions or formulations, at the same time, wherein one composition includes the *p53* expression vector and the other includes the DNA damaging agent.

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Alternatively, the p53 treatment may precede or follow the DNA damaging agent treatment by intervals ranging from minutes to weeks. In embodiments where the DNA damaging factor and p53 expression vector are applied separately to the cell, one would generally ensure that a significant period of time did not expire between the time of each

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delivery, such that the DNA damaging agent and expression vector would still be able to exert an advantageously combined effect on the cell. In such instances, it is contemplated that one would contact the cell with both agents within about 6 hours to one week of each other and, more preferably, within about 24-72 hours of each other, with a delay time of only about 48 hours being most preferred. In some situations, it may be desirable to extend the time period for treatment significantly, however, where several days (2, 3, 4, 5, 6 or 7) to several weeks (1, 2, 3, 4, 5, 6, 7 or 8) lapse between the respective administrations.

It also is conceivable that more than one administration of either the *p53* construct or the DNA damaging agent will be desired. Various combinations may be employed, where *p53* is "A" and the DNA damaging agent is "B":

A/B/A	B/A/B	B/B/A	A/A/B	B/A/A	A/B/B	A/B/B/B
B/B/B/A	B/B/A/B	A/A/B/B	A/B/A/B	A/B/B/A	B/B/A/A	B/A/B/B
B/A/B/A	B/A/A/B	A/A/A/B	B/A/A/A	A/B/A/A	A/A/B/A	B/A/A/B

To achieve cell killing, both agents are delivered to a cell in a combined amount effective to kill the cell.

DNA damaging agents or factors are defined herein as any chemical compound or treatment method that induces DNA damage when applied to a cell. Such agents and factors include radiation and waves that induce DNA damage such as,  $\gamma$ -irradiation, X-rays, UV-irradiation, microwaves, electronic emissions, and the like. A variety of chemical compounds, also described as "chemotherapeutic agents", function to induce DNA damage, all of which are intended to be of use in the combined treatment methods disclosed herein. Chemotherapeutic agents contemplated to be of use, include, *e.g.*, adriamycin, 5-fluorouracil (5FU), etoposide (VP-16), camptothecin, actinomycin-D, mitomycin C, cisplatin (CDDP) and even hydrogen peroxide. The invention also encompasses the use of a combination of one or more DNA damaging agents, whether radiation-based or actual compounds, such as the use of

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X-rays with cisplatin or the use of cisplatin with etoposide. In certain embodiments, the use of cisplatin in combination with a p53 expression vector is particularly preferred.

In treating cancer according to the invention, one would contact the tumor cells with a DNA damaging agent in addition to the expression vector. This may be achieved by irradiating the localized tumor site with DNA damaging radiation such as X-rays, UV-light, g-rays or even microwaves. Alternatively, the tumor cells may be contacted with the DNA damaging agent by administering to the subject a therapeutically effective amount of a pharmaceutical composition comprising a DNA damaging compound such as, adriamycin, 5-fluorouracil, etoposide, camptothecin, actinomycin-D, mitomycin C, or more preferably, cisplatin. The DNA damaging agent may be prepared and used as a combined therapeutic composition, or kit, by combining it with a p53 expression vector, as described above.

Agents that directly cross-link polynucleotides, specifically DNA, are envisaged and are shown herein, to eventuate DNA damage leading to a synergistic antineoplastic combination. Agents such as cisplatin, and other DNA alkylating may be used. Cisplatin has been widely used to treat cancer, with efficacious doses used in clinical applications of 20 mg/m² for 5 days every three weeks for a total of three courses. Cisplatin is not absorbed orally and must therefore be delivered via injection intravenously, subcutaneously, intratumorally or intraperitoneally.

Agents that damage DNA also include compounds that interfere with DNA replication, mitosis and chromosomal segregation. Such chemotherapeutic compounds include adriamycin, also known as doxorubicin, etoposide, verapamil, podophyllotoxin, and the like. Widely used in a clinical setting for the treatment of neoplasms, these compounds are administered through bolus injections intravenously at doses ranging from 25-75 mg/m² at 21 day intervals for adriamycin, to 35-50 mg/m² for etoposide intravenously or double the intravenous dose orally.

Agents that disrupt the synthesis and fidelity of polynucleotide precursors and subunits also lead to DNA damage. As such a number of polynucleotide precursors have been developed.

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Particularly useful are agents that have undergone extensive testing and are readily available. As such, agents such as 5-fluorouracil (5-FU), are preferentially used by neoplastic tissue, making this agent particularly useful for targeting to neoplastic cells. Although quite toxic, 5-FU, is applicable in a wide range of carriers, including topical, however intravenous administration with doses ranging from 3 to 15 mg/kg/day being commonly used.

Other factors that cause DNA damage and have been used extensively include what are commonly known as  $\gamma$ -rays, X-rays, and/or the directed delivery of radioisotopes to tumor cells. Other forms of DNA damaging factors are also contemplated such as microwaves and UV-irradiation. It is most likely that all of these factors effect a broad range of DNA damage, or the precursors of DNA, the replication and repair of DNA, and the assembly and maintenance of chromosomes. Dosage ranges for X-rays range from daily doses of 50 to 200 roentgens for prolonged periods of time (3 to 4 weeks), to single doses of 2000 to 6000 roentgens. Dosage ranges for radioisotopes vary widely, and depend on the half-life of the isotope, the strength and type of radiation emitted, and the uptake by the neoplastic cells.

The skilled artisan is directed to "Remington's Pharmaceutical Sciences" 15th Edition, chapter 33, in particular pages 624-652. Some variation in dosage will necessarily occur depending on the condition of the subject being treated. The person responsible for administration will, in any event, determine the appropriate dose for the individual subject. Moreover, for human administration, preparations should meet sterility, pyrogenicity, general safety and purity standards as required by FDA Office of Biologics standards.

The inventor proposes that the regional delivery of p53 expression vectors to patients with p53-linked cancers will be a very efficient method for delivering a therapeutically effective gene to counteract the clinical disease. Similarly, the chemo- or radiotherapy may be directed to a particular, affected region of the subject's body. Alternatively, systemic delivery of the expression vector or the DNA damaging agent may be appropriate in certain circumstances, for example, where extensive metastasis has occurred.

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Cytokine therapy also has proven to be an effective partner for combined therapeutic regimens. Various cytokines may be employed in such combined approaches. Examples of cytokines include IL-1α IL-1β, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, TGF-β, GM-CSF, M-CSF, G-CSF, TNFα, TNFβ, LAF, TCGF, BCGF, TRF, BAF, BDG, MP, LIF, OSM, TMF, PDGF, IFN-α, IFN-β, IFN-γ. Cytokines are administered according to standard regimens, as described below, consistent with clinical indications such as the condition of the patient and relative toxicity of the cytokine.

In addition to combining *p53*-targeted therapies with chemo-, radio and cytokine therapies, it also is contemplated that combination with other gene therapies will be advantageous. For example, targeting of K-ras and p53 mutations at the same time may produce an improved anti-cancer treatment. Any other tumor-related gene conceivably can be targeted in this manner, for example, p21, p16, p27, E<sub>2</sub>F, Dp family of genes, Rb, APC, DCC, NF-1, NF-2, WT-1, MEN-II, BRCA1, VHL, FCC, MCC, other ras molecules, myc, neu, raf, erb, src, fms, jun, trk, ret, gsp, hst, bcl, and abl. It also may be desirable to combine p53 therapy with an antibody-based gene therapy treatment involving the use of a single-chain antibody construct in which the antibody binds to any of the foregoing molecules, or in combination with genes encoding one or more of the cytokines listed above. It also may be advantageous to combine p53 with other genes that have been implicated in apoptotic processes, e.g., adenovirus E1A, Bax, Bcl-X<sub>S</sub>, etc.

# F. Kits

All the essential materials and reagents required for inhibiting tumor cell proliferation may be assembled together in a kit. This generally will comprise selected adenoviral expression vectors. Also included may be various media for replication of the expression vectors and host cells for such replication. Such kits will comprise distinct containers for each individual reagent.

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When the components of the kit are provided in one or more liquid solutions, the liquid solution preferably is an aqueous solution, with a sterile aqueous solution being particularly preferred. For *in vivo* use, the expression vector may be formulated into a pharmaceutically acceptable syringeable composition. In this case, the container means may itself be an inhalent, syringe, pipette, eye dropper, or other such like apparatus, from which the formulation may be applied to an infected area of the body, such as the lungs, injected into an animal, or even applied to and mixed with the other components of the kit.

The components of the kit may also be provided in dried or lyophilized forms. When reagents or components are provided as a dried form, reconstitution generally is by the addition of a suitable solvent. It is envisioned that the solvent also may be provided in another container means.

The kits of the present invention also will typically include a means for containing the vials in close confinement for commercial sale such as, e.g., injection or blow-molded plastic containers into which the desired vials are retained. Irrespective of the number or type of containers, the kits of the invention also may comprise, or be packaged with, an instrument for assisting with the injection/administration or placement of the ultimate complex composition within the body of an animal. Such an instrument may be an inhalent, syringe, pipette, forceps, measured spoon, eye dropper or any such medically approved delivery vehicle.

# G. Animal Model for Microscopic Tumor Seeding and Residual Disease

Another aspect of the present invention involves the development of an animal model for the analysis of microscopic residual carcinomas and microscopic seeding of body cavities. "Carcinoma," as used herein, may refer to a single cell or a multicellular tumor mass. In microscopic disease, the "tumor" will consist or one or a few carcinoma cells which cannot be observed with the naked eye.

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The animal model described herein is particularly advantageous mimicking (i) the post surgical environment of head and neck cancer patients, particularly in advanced stages of disease and (ii) the body cavity of an affected subject wherein microscopic carcinoma has been established. The model, similar to other animal models for cancer, derives from inoculation of tumor cells into an animal. A distinction, however, lies in the creation, subcutaneously, of a pouch that is a physiologic equivalent of a natural body cavity or a post-surgical cavity created by the excision of a tumor mass.

The instant inventions exemplifies nude mice as the model organism. Virtually any animal may be employed, however, for use according to the present invention. Particularly preferred animals will be small mammals that are routinely used in laboratory protocols. Even more preferred animals will be those of the rodent group, such as mice, rats, guinea pigs and hamsters. Rabbits also are a preferred species. The criteria for choosing an animal will be largely dependent upon the particular preference of an investigator.

The first step is to create a tissue flap in the experimental animal. The term "tissue flap" means any incision in the flesh of the animal that exposes the target tissue. It is generally preferred that an incision be made in the dorsal flank of an animal, as this represents a readily accessible site. However, it will be understood that an incision could well be made at other points on the animal, and the choice of tissue sites may be dependent upon various factors such as the particular type of therapeutics that are being investigated.

Once a target tissue site is exposed, carcinoma cells, either individually or in microscopic tumors, are contacted with the tissue site. The most convenient manner for seeding the cancer cells into the tissue site is to apply a suspension of tissue culture media containing the cells to the exposed tissue. Cancer cell application may be achieved simply using a sterile pipette or any other convenient applicator. Naturally, this procedure will be conducted under sterile conditions.

In one embodiment,  $2.5 \times 10^6$  SCCHN cells are inoculated into the exposed tissue flap of a nude mouse. Those of skill in the art will be able to readily determine, for a given purpose, what the appropriate number of cells will be. The number of cells will be dependent upon various factors, such as the size of the animal, the site of incision, the replicative capacity of the tumor cells themselves, the time intended for tumor growth, the potential anti-tumor therapeutic to be tested, and the like. Although establishing an optimal model system for any particular type of tumor may require a certain adjustment in the number of cells administered, this in no way represents an undue amount of experimentation. Those skilled in the area of animal testing will appreciate that such optimization is required.

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This can be accomplished, for example, by conducting preliminary studies in which differing numbers of cells are delivered to the animal and the cell growth is monitored following resealing of the tissue flap. Naturally, administering larger numbers of cells will result in a larger population of microscopic residual tumor cells.

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In the present study the flaps were effectively sealed using mattress sutures. However, it is envisioned that persons skilled in the art may use any of a variety of methods routinely used to seal the incision such as the use of adhesives, clamps, stitches, sutures, *etc.*, depending on the particular use contemplated.

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# H. Examples

The following examples are provided merely to illustrate certain specific embodiments and should not be considered as limiting the scope of the invention in any way.

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#### **EXAMPLE 1**

# Growth Suppression of Human Head and Neck Cancer Cells by the Introduction of a Wild-Type *p53* Gene *via* a Recombinant Adenovirus

# **Materials and Methods**

Cell Lines and Culture Conditions. Human SCCHN cell lines Tu-138 and Tu-177 were both established at the Department of Head and Neck Surgery, M.D. Anderson Cancer Center. Tu-138 and Tu-177 were established from a gingivo-labial moderately differentiated squamous carcinoma and a poorly differentiated squamous carcinoma of the larynx, respectively. Both cell lines were developed via primary explant technique and are cytokeratin positive and tumorigenic in athymic nude and SCID mice. These cells were grown in DMEM/F12 medium supplemented with 10% heat-inactivated fetal bovine serum (FBS) with penicillin/streptomycin.

Recombinant Adenovirus Preparation and Infection. The recombinant *p53* adenovirus (Ad5CMV-*p53*) (Zhang *et al.*, 1994) contains the cytomegalovirus (CMV) promoter, wild-type *p53* cDNA, and SV40 polyadenylation signal in a mini-gene cassette inserted into the E1-deleted region of modified adenovirus type 5 (Ad5). Viral stocks were propagated in 293 cells. Cells were harvested 36-40h after infection, pelleted, resuspended in phosphate-buffered saline, lysed, and cell debris was removed by subjecting to CsCl gradient purification. Concentrated virus was dialyzed, aliquoted and stored at -80°C. Infection was carried out by addition of the virus in DMEM/F12 medium and 2% FBS to cell monolayers, the cells were incubated at 37°C for 60 min with constant agitation, then complete medium (DMEM/F12/10% FBS) was added and the cells were incubated at 37°C for the desired length of time.

Northern Blot Analysis. Total RNA was isolated by the acid-guanidinium thiocyanate method of (Chomczynski and Sacchi, 1987). Northern analyses were performed on 20  $\mu$ g of total RNA. The membrane was hybridizated with a p53 cDNA probe labeled by the random primer method in 5 x SSC/5X Denhardt's solution/0.5% SDS/denatured salmon sperm DNA (20  $\mu$ g/ml). The membrane was also stripped and reprobed with GAPDH cDNA for RNA loading

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control. The relative quantities of p53 expressed were determined by densitometer (Molecular Dynamics Inc., Sunnyvale, CA).

Western Blot Analysis. Total cell lysates were prepared by sonicating the cells 24-h post-infection in RIPA buffer (150 mM NaCl, 1.0% NP-40, 0.5% DOC, 0.1% SDS, 50 mM Tris, pH 8.0) for 5s. Fifty micrograms of protein from samples were subjected to 10% SDS-PAGE and transferred to Hybond-ECL membrane (Amersham). The membrane was blocked with Blotto/Tween (5% nonfat dry milk, 0.2% Tween 20, 0.02% sodium azide in phosphate-buffered saline) and probed with the primary antibodies, mouse anti-human *p53* monoclonal antibody PAb1801 and mouse anti-human β-actin monoclonal antibody (Amersham), and the secondary antibody, horseradish peroxidase-conjugated goat anti-mouse IgG (Boehringer Mannheim, Indianapolis, IN). The membrane was processed and developed as the manufacturer suggested.

Immunohistochemical Analysis. The infected cell monolayers were fixed with 3.8% formalin and treated with 3%  $H_2O_2$  in methanol for 5 min. Immunohistochemical staining was performed by using the Vectastain Elite kit (Vector, Burlingame, CA). The primary antibody used was the anti-p53 antibody PAb1801, and the secondary antibody was an avidin-labeled anti-mouse IgG (Vector). The biotinylated horseradish peroxidase ABC complex reagent was used to detect the antigen-antibody complex. Preadsorption controls were used in each immunostaining experiment. The cells were then counterstained with Harris hematoxylin (Sigma Chemical Co., St. Louis, MO).

Cell Growth Assay. Cells were plated at a density of 2x10<sup>4</sup> cells/ml in 6-well plates in triplicate. Cells were infected with either wild-type (Ad5CMV-p53) or replication-deficient adenovirus as a control. Cells were harvested every 2 days, counted and their viability was determined by trypan blue exclusion.

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Inhibition of Tumor Growth In Vivo. The effect of Ad5CMV-p53 on established subcutaneous tumor nodules was determined in nude mice in a defined pathogen free environment. Experiments were reviewed and approved by institutional committees for both animal care and utilization and for recombinant DNA research. Briefly, following induction of acepromazine/ketamine anesthesia, three separate subcutaneous flaps were elevated on each animal and 5 x 10<sup>6</sup> cells in 150 ml of complete media were injected subcutaneously into each flap using a blunt needle; the cells were kept in the pocket with a horizontal mattress suture. Four animals were used for each cell line. After 4 days, the animals were re-anesthetized and the flaps were re-elevated for the delivery of 100 ml of 1) Ad5CMV-p53 (50 MOI) in the right anterior flap; 2) replication-defective virus (50 MOI) in the right posterior flap; and 3) transport medium alone, in the left posterior flank. All injection sites had developed subcutaneous visual and palpable nodules before treatment was administered. Animals were observed daily and sacrificed on day 20. in vivo tumor volume was calculated by assuming a spherical shape with the average tumor diameter calculated as the square root of the product of cross-sectional diameters. Following sacrifice, excised tumors were measured 3-dimensionally by microcalipers to determine tumor volume. A non-parametric Friedman's 2-way ANOVA test was used to test the significance of the difference between means of samples; the SPSS/PC+ software package (SPSS Inc., Chicago, II) was used.

# **Results**

Adenoviral Infection of SCCHN Cells. The conditions for optimal adenoviral transduction of Tu-138 and Tu-177 cells were determined by infecting these cells with adenovirus expressing the E. coli  $\beta$ -gal gene. The transduction efficiency was assessed by counting the number of blue cells after X-gal staining. There appeared to be a linear relationship between the number of infected cells and the number of adenovirus particles used. Cells inoculated with a single dose of 100 MOI  $\beta$ -gal adenovirus exhibited 60% blue cells (FIG. 1), and this was improved to 100% by multiple infections. The transduction efficiency of this vector in SCCHN cells is quite different from that of other cell lines previously examined: HeLa,

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HepG2, LM2 and human non-small cell lung cancer cell lines showed 97% to 100% infection efficiencies after incubation with 30 to 50 MOI  $\beta$ -gal adenovirus (Zhang *et al.*, 1994).

Expression of Exogenous p53 mRNA in Adenovirus Infected SCCHN Cells. Two human SCCHN cell lines were chosen for this study: both cell lines Tu-138 and Tu-177 possess The recently created recombinant wild-type p53 adenovirus, a mutated p53 gene. Ad5CMV-p53, was used to infect Tu-138 and Tu-177 cells. Twenty-four hours after infection, total RNA was isolated and northern blot analysis was performed. The transformed primary human embryonal kidney cell line 293 was used as a positive control because of its high level of expression of the p53 gene product, whereas K562, a lymphoblastoma cell line with a homozygous deletion of the p53 gene, was the negative control. The levels of the 2.8-kb endogenous p53 mRNA detected in the samples isolated from mock-infected cells and from the cells infected with a replication-defective adenovirus, dl312 were similar. Up to 10-fold higher levels of exogenous 1.9-Kb p53 mRNA were present in the cells infected with Ad5CMV-p53, indicating that the exogenous p53 cDNA was successfully transduced into these cells and efficiently transcribed. Interestingly, the level of endogenous p53 mRNA in these cells was 5-fold higher than in the experimental controls. Northern blots exhibited no evidence of Ad5CMV-p53 (DNA) contamination of RNA.

Expression of p53 Protein in Adenovirus-Infected SCCHN Cells. Western blot analysis was performed to compare the levels of p53 mRNA to the amount of p53 protein produced. A p53 band, recognized by monospecific anti-p53 antibody, PAb1801, was observed in cellular extracts isolated from all samples except K562 cells. Cell line 293 showed high levels of p53 protein. Samples isolated from mock-infected Tu-138 and Tu-177 cells exhibited low levels of p53 protein. The level of p53 expression remained similar in those cells infected with the dl312 adenovirus. The levels of p53 antigen detected in Ad5CMV-p53-infected cells were significantly higher than the levels of the endogenous mutated proteins in both cell lines. This result demonstrates that the exogenous p53 mRNA produced from cells infected with Ad5CMV-p53 is efficiently translated into immunoreactive p53 protein. Furthermore,

immunohistochemical analysis of cells infected with Ad5CMV-p53 revealed the characteristic nuclear staining of p53 protein, while mock-infected cells failed to show similar staining despite the presence of the p53 protein in these cells. This inability to detect the protein may be attributable to the insensitivity of the assay.

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Effect of Exogenous p53 on SCCHN Cell Growth In Vitro. Cells infected with control virus dl312 had growth rates similar to those of the mock-infected cells (FIG. 2A and 2B), whereas, growth of the Ad5CMV-p53-infected Tu-138 (FIG. 2A) and Tu-177 (FIG. 2B) cells was greatly suppressed. Twenty-four hours after infection, an apparent morphologic change occurred, with portions of the cell population rounding up and their outer membranes forming blebs. These are part of a series of histologically predictable events that constitute programmed cell death. The effect was more prominent for Tu-138 than for Tu-177 cells. Cells infected with the replication-defective adenovirus, dl312, demonstrated normal growth characteristics with no histomorphologic abnormalities. Growth assays were reproducible in four repeated experiments.

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Inhibition of Tumor Growth *In Vivo*. Four animals were tested for each cell lines. One animal in the Tu-177 group died following the second flap surgery and delivery of the therapeutic interventions, presumably due to profound anesthesia and subsequent mutilation by cage mates. Necropsy revealed no evidence of metastasis or systemic effects. Sizable tumors are apparent on both posterior flaps of the animals (i.e., the sites that did not receive Ad5CMV-p53). The lack of tumor progression is significant in the right anterior flaps of the animals, which received Ad5CMV-p53 (p < .04) That Tu-177 cells have a slower growth rate has previously been established in these animals. Two animals in the Tu-138 group were killed early because they were experiencing rapid growth and ulceration of the control tumor sites. All surgical sites had developed lesions of at least 9 mm<sup>3</sup> before intervention. The tumor volumes on necropsy are shown in Table 4. Differences in volume were not statistically significant in the Tu-177 group which may be a reflection of the limited sample size.

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TABLE 4

Effect of Ad5CMV-p53 on Tumor Growth in Nude Mice<sup>a</sup>

Treatment	Mean v [mm <sup>3</sup> ±			
	TU-138(4)	Tu-177(3)		
Ad5CMV-p53	$36.0 \pm 23.5$	19 ± 14.9		
Ad5(dl312)	$1187.5 \pm 419.0$	$226 \pm 84.0$		
Medium	$1909.3 \pm 776.8$	$454 \pm 276.8$		
Significance	p value	p value		
<i>p53</i> <sup>b</sup> : dl312	0.04	0.09		
p53: Medium	0.05	0.09		

<sup>&</sup>lt;sup>a</sup>The cells were injected subcutaneously at  $5 \times 10^6$  cells/flap. Tumor sizes were determined at day 20 after treatment. Numbers in parentheses represent the number of animals evaluated. <sup>b</sup>Ad5CMV-*p53* is abbreviated as *p53*; dl312 as an abbreviation for Ad5(dl312).

# EXAMPLE 2

in vivo Molecular Therapy with p53 Adenovirus forMicroscopic Residual Head and Neck Squamous Carcinoma

# **Materials and Methods**

Cell Lines and Culture Conditions. Human SCCHN cell lines Tu-138, Tu-177, MDA 686-LN, and MDA 886 were all established have been previously characterized (Clayman *et al.*, 1993; Sacks *et al.*, 1988). These cells were grown in Dulbecco's modified Eagle's medium (DMEM/F12) supplemented with 10% heat-inactivated fetal bovine serum (FBS) and penicillin/streptomycin.

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Recombinant Adenovirus Preparation and Infection; Cell Growth Assay; Western Blot Analysis. All the procedures have been previously described in Example 1. Cell growth assays were all performed in triplicate.

In Vivo Transduction with β-Galactosidase Adenovirus. X-gal staining of tissue specimens were performed on O.C.T. frozen tissue sections to determine transduction efficiency. Eight-micrometer thick specimens were washed in cold PBS and fixed in 0.5% glutaraldehyde at room temperature for 5 minutes. Slides were then washed twice with 4°C PBS and incubated 4 h in X-gal solution (1.3 mM MgCl<sub>2</sub>; 15 mM, NaCl; 44 mM Hepes Buffer pH7.4; 3 mM Potassium ferricyanide; 3 mM Potassium ferrocyanide; 2% X-gal in DMF). Slides were counterstained with hematoxylin and eosin.

Immunohistochemical Analysis. Formalin-fixed paraffin-embedded in vivo animal experimental tissues were cut at 4-5 µm, dried at 60°C, deparaffinized, and hydrated with distilled water. Sections were then treated with 0.5% saponin in distilled water and rinsed in several changes of distilled water; endogenous peroxidase activity was blocked with 3% hydrogen peroxide in methanol, followed by rinsing in several changes of distilled water. Sections were microwave-irradiated in distilled water for 3 min using a Sharp Model R9H81 microwave oven operating at a frequency of 2450 MHz at 700 watts. After cooling, sections were washed in several changes of distilled water and placed in PBS; immunochemical studies were performed by using the avidin-biotin-peroxidase complex (ABC) method of Hsu et al., (1981) in the following manner: sections were blocked with normal horse serum and incubated overnight at 4°C with rabbit anti-human p53 polyclonal antibody, clone OM-1, 1:80 (Signet Laboratories, Denham, MA). An anti-rabbit IgG Elite kit (Vector Laboratories, Burlingame, CA) was then used to apply biotinylated anti-rabbit IgG and ABC complexes, which were incubated for 45 min each. The immunostaining reaction was visualized by using 0.5% DAB in PBS containing 0.01% hydrogen peroxide (pH 7.6), counterstained with 0.01% toluidine blue, dehydrated, cleared, and mounted in Permount. To verify the specificity of the immunostaining reaction, immunoperoxidase staining was performed, using the same method as on test samples,

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on a known positive cytospin of a tissue culture of a squamous carcinoma cell line as well as on a negative rabbit monoclonal antibody control.

Inhibition of Tumor Growth *In Vivo*. This procedure was performed as described in Example 1. All surgical sites were evaluated pathologically as well as by necropsy analysis for systemic toxicity.

# Results

Effect of Exogenous p53 on SCCHN Cell Growth In Vitro. Example 1 described the in vitro inhibition of cell growth by Ad5CMV-p53 in SCCHN cell lines with endogenously mutated p53. This present example is to determine whether SCCHN cell lines with endogenous wild-type p53 would be similarly affected. The effect of Ad5CMV-p53 on nonmalignant fibroblasts is also investigated.

Four human SCCHN cell lines were chosen for this study: Tu-138 and Tu-177 possess a mutated *p53* gene, whereas MDA 686-LN and 886 both are homozygous for the wild-type *p53* gene. A fibroblast cell line derived from normal fibroblast outgrowth, which is karyotypically normal and nontumorigenic, was used as a nonmalignant control cell line. Cells infected with the control virus, dl312, had growth rates similar to those of the mock-infected cells, whereas the growth of tumor cells infected with Ad5CMV-*p53* was significantly suppressed (FIG. 3A, FIG. 3B, FIG. 3C AND FIG. 3D). Twenty-four hours to forty-eight hours after infection, an apparent morphologic change occurred in all tumor cells, with portions of the cell population rounding up and their outer membranes forming blebs. These are part of a series of histologically predictable events that constitute programmed cell death. The effect occurred earlier in cells with endogenous mutated *p53* than in those with wild-type *p53*. Cells infected with replication-defective adenovirus dl312 demonstrated normal growth characteristics with no histomorphologic abnormalities. Growth assays were reproducible in four repeated experiments.

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Expression of Exogenous p53 Protein in Adenovirus Infected Normal Fibroblasts and its Effect on Growth Rate. Additionally, the effect of the Ad5CMV-p53 on karyotypically normal and non-tumorigenic fibroblast cell lines was also investigated. These cells were isolated during the establishment of primary tumor cell lines. Twenty-four hours after infection, Western blot analysis was performed to compare the levels of protein produced by the different infected cell types. A p53 band, recognized by the monospecific anti-p53 antibody, PAb1801, was observed in cellular extracts isolated from all samples infected with the Ad5CMV-p53. As has been in Example 1, cell line Tu-138 infected with the p53 adenovirus showed high levels of p53 protein following transduction and served as a control. The level of p53 expression remained similar in both mock-infected and dl312 infected cells. The Ad5CMV-p53 infected fibroblasts showed higher levels of p53 protein than that of the control cells. This result indicates that the p53 gene is efficiently translated into normal fibroblasts infected with Ad5CMV-p53 as evidenced by production of immunoreactive p53 protein. The protein expression and transduction fibroblasts Ad5CMV-p53 infected were verified of efficiency of cytospins This fibroblast cell line exhibited normal growth rate and immunohistochemical analysis. intervention (mock, replication-defective virus, independent of the morphology, Ad5CMV-p53) (FIG. 4). These experiments were repeated twice and also verified in other normal human fibroblast cell lines.

In Vivo Transduction Efficiency. To measure the efficiency of gene transfer in vivo, the subcutaneous flap site was resected 72 hours following the molecular or control intervention. Dose-response experiments with the adenovirus  $\beta$ -galactosidase marker vector demonstrate dose-response transduction efficiency in this model. This was confirmed with immunohistochemical analysis 4 days following infection with Ad5CMV-p53. Both groups of experiments exhibited an *in vivo* dose response which had been previously described *in vitro* (Example 1). In no instances did doses of virus exceeding  $10^{10}$  PFU effect expression of p53 in other organ systems, including brain, liver, lung, heart, abdominal visceral organs, and skin. These experiments illustrated a dose-response relationship between viral titer and transduction

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efficiency as well as the possibility of achieving extensive transient expression of the transduced gene within the desired surgical model field.

Suppression of Tumor Growth In Vivo. Studies were designed to determine whether in vivo Ad5CMV-p53 mediated gene transfer would affect the establishment or growth of SCCHN cells implanted into a subcutaneous flap. To achieve this aim a microscopic residual disease model was created. In this model, three subcutaneous flaps were elevated on athymic nude female mice and 2.5x10<sup>6</sup> of tumor cells were seeded by pipetting. Instead of allowing the tumor cells to form nodules (generally occurring in 4 days), a single dose of molecular intervention at 48 hours following tumor cells seeding. In this manner, although no gross tumors were present, microscopic tumor cells were within the surgical site mimicking the clinical dilemma of surgical excision of all gross tumor. The development of tumors was directly related to the number of tumor cells, the time allotted for implantation, and the dose of Ad5CMV-p53. Of the mice which received microscopically implanted tumor cells (2.5 x 10<sup>6)</sup> and were treated with Ad5CMV-p53 at 10<sup>8</sup> plaque forming units (PFU) or greater, only two developed tumors, both implanted with the wild-type p53 cell line (MDA 886-LN). All other cell lines exhibited absence of tumor development (Table 5). These experiments clearly demonstrate that the growth of microscopic tumor cells can be effectively suppressed in vivo if exposed to the Ad5CMV-p53. Tumor formation was evaluated at the end of a 12 week period (earlier animal sacrifice in circumstances of excessive tumor burden) by gross and histologic analysis of the surgical sites. The data of tumor establishment is summarized in Table 5.

TABLE 5

Effect of Ad5CMV-p53 on Tumorigenicity in a
Microscopic Residual Disease Model of SCCHN

**Cell Line** 

# Treatment

# No. Mice Developing Tumors/Total Mice

	Vehicle								
	PBS	dl312	Ad5CMV-p53						
Tu-138 (homozygous mutation p53)	8/8	8/8	0/8						
Tu-177 (homozygous mutation p53)	8/8	8/8	0/8						
686-LN (homozygous wild-type p53)	5/8	5/8	0/8						
886 (homozygous wild-type <i>p53</i> )	6/6	6/6	2/6						

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Immunohistochemical analysis was performed on the tumor sections of experimental animals. This cell line possesses the wild-type endogenous *p53* gene. There was a lack of significant basal immunostaining with viable tumor of MDA 686-LN (mock-infection). 10<sup>7</sup> PFU Ad5CMV-*p53* exhibits peripheral tumor necrosis with immunostaining in the more central portion of the tumor. 10<sup>8</sup> PFU Ad5CMV-*p53* reveals total necrosis of the tumor with immunostaining found in the entire surgical pocket with multiple layers expressing protein including stroma and superficial muscular layers. 10<sup>9</sup> PFU Ad5CMV-*p53*) shows similar results to that of 10<sup>8</sup> PFU Ad5CMV-*p53*, however increased exogenous *p53* expression throughout the surgical site and edema are prominent.

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Using animals, which served as their own internal controls, implants of  $4.0 \times 10^6$  or more cells significantly increased the establishment of subcutaneous implants as compared to the tumor implantation of  $2.5 \times 10^6$  cells (P < 0.01), even when treated at the surgical site with Ad5CMV-p53 48 h after inoculation. Allowing implanted cells to establish for 72 or 96 h prior to the Ad5CMV-p53 intervention similarly increased tumor take. Dose-response experiments established that  $10^8$  and  $10^9$  PFU of the Ad5CMV-p53 were equally effective in inhibiting tumor burdens of  $2.5 \times 10^6$  cells implanted for 48 h (FIG. 6). Endogenous p53 status of implanted

tumor cell lines (whether homozygous mutated or wild-type *p53*) had little impact on the effectiveness of the Ad5CMV-*p53* in the cessation of tumor development.

#### EXAMPLE 3

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# Apoptosis Induction Mediated by Wild-Type *p53* Adenovirus Gene Transfer in Squamous Cell Carcinoma of Head and Neck

#### **Materials and Methods**

DNA was analyzed in a 1% agarose gel.

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Cell Lines and Culture Conditions; Recombinant Adenovirus Preparation and Infection. All procedures were performed and cell lines maintained as previously described in Examples 1 and 2.

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DNA Fragmentation Analysis. Following incubation with wild-type *p53* adenovirus as well as replication defective adenovirus controls at various time intervals, cells were harvested, resuspended in 300 μl of PBS with the addition of 3 ml of extraction buffer (10 mM Tris, pH 8.0, 0.1M EDTA, 20 μg/ml RNAse, 0.5% SDS) and incubated at 37°C for 1-2 h. At the end of incubation, proteinase K was added to a final concentration of 100 μg/ml and the solution placed in a 50°C water bath for at least 3 h. DNA was extracted once with equal volume of 0.5 M Tris (pH 8.0) saturated phenol then the extraction repeated with phenol/chloroform. Precipitated

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Cell Fixation. For TUNEL method, the cells were fixed in 1% formaldehyde in PBS (pH 7.4) for 30 min on ice. Cells were then washed with 3 ml of PBS, resuspended in 70% ice-cold ethanol and stored at -20°C until used. For cell-cycle analysis, cells were fixed in 70% ice-cold ethanol only.

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Terminal Deoxynucleotidyl Transferase Assay. The assay was performed according to Gorczyca et al., procedure (Gorczyca et al., 1993). Briefly, after fixation and washing, cells

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were resuspended in 50  $\mu$ l of TdT buffer containing 0.2 M sodium cacodylate (pH 7.0), 2.5 mM Tris-HCl, 2.5 mM  $C_0Cl_2$  (Sigma Chemical Company, St. Louis, MO), 0.1 mM DTT (Sigma Chemical Company), 0.25 mg/ml BSA (Sigma Chemical Company), 5 units of terminal transferase (Boehringer Mannheim Biochemicals, Indianapolis, IN), and 0.5 nmoles biotin-16-dUTP along with dATP,dGTP and dCTP at a concentration of 20  $\mu$ M. Controls were prepared by incubating a separate aliquot of each test sample without d-UTP. The cells were incubated in the solution at 37 $^{0}$ C for 30 min, rinsed in PBS, and resuspended in 100  $\mu$ l of, FITC, the staining solution containing 4X SSC, 0.1% Triton X-100 and 2.5  $\mu$ g/ml fluoresceined avidin (Vector Labs. Inc., Burlingame, CA). Tubes were incubated for 30 min in the dark at room temperature. Cells were rinsed in PBS with 0.1% Triton X-100 and resuspended in 0.5 ml PBS containing propidium iodide (5  $\mu$ g/ml) and 70  $\mu$ l (1 mg/ml) RNAse. Tubes were incubated in the dark on ice for 30 min prior to flow cytometric analysis.

Flow Cytometry Analysis. All samples were analyzed using an EPICS Profile II flow cytometer (Coulter Corp., Hialeah, FL) with the standard optical configuration. At least 5,000 events were collected for each sample. Positivity for TdT end-labeling was determined by subtracting the control histogram from the test histogram using the immuno-4 program of the Elite workstation software (Coulter Corp., Hialeah, FL).

Cell Growth Assay. Cells were plated and growth was monitored as described in Example 1.

In Vivo Analysis for Apoptosis. Gene therapy in a microscopic residual disease model of SCCHN has been described above in Example 2.

In Situ End-labeling. The procedure was performed as previously described (Wijsman et al., 1993). Briefly, paraffin sections were dewaxed in xylene for 5 min three times each, and progressively hydrated by immersing the slides for 3 min each in 100%, 90%, 70% and 30% ethanol solutions. Endogenous peroxidase was inactivated by immersing the slides for 20 min in

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 $0.75\%~H_2O_2~v/v$  in 100% methanol. After the slides were washed in PBS, sections were digested with 0.1%~pepsin (Fisher Scientific, Houston, TX) w/v in 0.1~N~HCl for 5 min at 37°C and extensively washed in PBS. Sections were then incubated in a moist chamber at 37°C for 1 h with an end-labeling cocktail including the following: terminal deoxynucleotidyl transferase 0.5~v unit/µl; biotinylated dUTP, 0.06~mM; 5X tdt buffer,  $10~\mu$ l; double-distilled water up to  $50~\mu$ l. The reaction was terminated by immersing the slides in a buffer containing 300 mM NaCl and 30 mM NaCitrate in double-distilled water. After the slides were washed in PBS, sections were incubated with horseradish peroxide-conjugated avidin for  $1~h~at~37^{\circ}C$  in a moist chamber. Staining was developed using 3,3'-diaminobenzidine and sections were counterstained with methyl green.

# **Results**

Growth suppression of SCCHN cell lines by p53 adenovirus. The Examples above demonstrate that the wild-type p53 gene can be efficiently transduced into SCCHN cell lines by a recombinant adenoviral vector. Consequently, the insulted tumor cells lose their ability to proliferate in vitro as well as in vivo. The suppression effect is independent of the endogenous p53 status of the cell lines. Previous growth rate analyses were carried out through a week period. The present example investigates the early effects of the wild-type p53 on SCCHN cells growth (i.e. earlier time intervals, hours).

Two representative cell lines were used in this study. Cell line, Tu-138, harbors a mutated *p53* gene whereas cell line, MDA 686LN, possesses a wild-type *p53* gene. Cells infected with replication-defective virus, dl312, had growth rates similar to those of the mock-infected cells (FIG. 5A and FIG. 5B). On the other hand, growth of the Ad5CMV-*p53* infected Tu138 (FIG. 5A) and MDA 686LN (FIG. 5B) cells were significantly suppressed. It appeared that the exogenous *p53* protein had an earlier and more profound growth suppression of the Tu138 as compared to the MDA 686LN. An apparent morphologic change was observed, with portions of the cell populations rounding up and their outer membranes forming blebs, resembling apoptosis, which occurred concomitantly with the initiation of the growth

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suppression. Cells infected with the replication-defective adenovirus, dl312, demonstrated normal growth characteristics with no histomorphologic abnormalities. Importantly, these effects were not observed following the p53 adenovirus infection of karyotypically normal fibroblasts, as detailed in Example 2 above, as well as human oral keratinocytes (immortalized but nontumorigenic).

DNA fragmentation analysis. One of the characteristic markers in apoptosis that distinguishes it from necrosis is the biochemically observable appearance of the ladder of DNA fragments. To confirm the notion that the cells had undergone apoptosis following the *p53* adenovirus infection, performed DNA fragmentation analysis was performed. Chromosomal DNA extracted from the viable cells following the replication-defective or the wild-type *p53* adenovirus infection were subjected to agarose gel electrophoresis. The appearance of DNA fragments equivalent to approximately 200 bp and their multiples was noticed in both cell lines. The fragmented DNA appeared at 22 h following the *p53* adenovirus infection in Tu-138 cell line whereas in the MDA 686LN cell line, fragmented DNA were visible at 30 h and more evident at 48 h following the wild-type *p53* adenovirus infection. No detectable fragmented DNA emerged from the mock-infected and dl312-infected cells.

In Vitro Terminal Deoxynucleotidyl Transferase Assay. Another characteristic marker of apoptosis is the morphological change and destruction of the structural organization of the nucleus which results in chromatin condensation. Electron microscopy has been used extensively to detect such ultra-structural alteration. However, recently flow cytometry methods for identifying apoptotic cells have gained favor due to the ability to scan and analyze cellular populations as compared to electron microscopy (Gorczyca et al., 1993). The inventor employed here the TUNEL (terminal deoxynucleotidyl transferase-mediated dUTP-biotin nick end-labeling) method (Gorczyca et al., 1993) which is based upon detection of the extensive DNA breakage to identify the apoptotic cells. Fifteen h following the p53 adenovirus infection, 4.4% of the viable Tu-138 cell population were in apoptotic stages as compared to none of the MDA 686LN (FIG. 6A and FIG. 6B). The number of apoptotic cells increased proportionally as the duration of

observation was also increased following p53 adenovirus incubation. Nearly 31% of the Tu-138 cells had undergone apoptosis at 22 h. Although delayed in initial induction of apoptosis, approximately 60% of the MDA 686LN cells were in apoptotic stages 48 h following the p53 adenovirus infection. Noteworthy, the percentage of apoptotic cells as determined by Tunel method may be significantly underestimated since only viable cells were subjected to the analysis. These data correlated well with the growth rate and DNA fragmentation analyses. There were no detectable cell populations undergoing apoptosis in control experiments using mock infection as well as replication defective virus controls (100 M.O.I.). Therefore, the apoptosis was not the function of transduced adenoviral gene products themselves.

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In Vivo Analysis for Apoptosis. The examples above show that the p53 adenovirus suppresses tumor formation in vivo. This example was designed to show whether the suppression of tumor growth in vivo was the consequence of apoptosis. In situ end-labeling analysis was performed to detect apoptotic cells in paraffin embedded sections obtained from example 2. Clearly, no staining was observed in the tissue sections isolated from MDA 686LN bearing animals which had received PBS treatment as a control. On the other hand, tissue sections isolated from MDA 686LN bearing mice treated with the wild-type p53 adenovirus showed highly positive staining, demonstrating that apoptosis indeed was the event involved in suppression of tumor growth in vivo.

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Further to these studies the inventor sought to determine whether the growth suppression is, in part, due to the cell cycle arrest by the induced p21 protein or primarily a result of apoptosis. Western blotting showed that the p21 protein was induced in the wild-type p53 adenovirus infected SCCHN cells. However, cell cycle analyses indicated that despite the elevated level of p21 protein in the p53 adenovirus infected cells, there was no significant accumulation of cells at the  $G_1$  stage when compared to S phase.

#### **EXAMPLE 4**

# Growth Suppression of Head and Neck Squamous Cells by *p53*-FLAG: an Effective Marker for Gene Therapy Trails

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#### Materials and Methods.

Cell Lines and Culture Conditions. Recombinant Adenovirus Preparation and Infection; Northern Blot Analysis; Western Blot Analysis; Cell Growth Assay; Immunohistochemical Staining in vitro Cell Layers. All procedures were performed and cell lines maintained as described in Example 1.

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Generation of the p53-FLAG Adenovirus. The p53 cDNA sequence was excised from the pC53-SN by digestion with BamH1 and cloned into the BamH1 site of pGEM7Z. A recombinant plasmid with the proper insert orientation was then digested with Accl and Kpn1 to remove 22 amino acids from the 3' end of p53 cDNA. A linker with Accl-Kpn1 compatible ends containing the sequence of the FLAG peptide including a stop codon was then ligated into the digested plasmid to create the p53-FLAG fusion gene. The resulting p53-FLAG fusion gene was then cloned into an expression vector with the human CMV promoter and SV polyadenylation signal. The final construct was subsequently inserted into a shuttle vector pXCJL.1 ((Zhang et al., 1994) to generate a recombinant p53-FLAG adenovirus.

In Vivo Microscopic Residual Disease Experiments. The studies were carried out in a defined pathogen free environment using the athymic nude model system described in Example 1. Two different sets of repeated experiments were performed. The first was a dose-response experiment using the AdCMV-p53-FLAG virus in three of the flaps at descending concentrations (10<sup>10</sup> pfu, 10<sup>9</sup> pfu, 10<sup>8</sup> pfu). The fourth flap served as a control and was randomized to either PBS or the replication defective adenovirus (DL312). The second study was performed using 10<sup>10</sup> pfu of AdCMV-p53-FLAG, AdCMV-p53, and replication defective adenovirus in three separate flaps. The fourth flap was inoculated with the same volume (100 μl) of sterile PBS.

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Forty eight hours after treatment, two of these animals were sacrificed and the flaps harvested for immunohistochemical analysis. The remaining animals were observed for 21 days and then sacrificed. Tumor volumes were measured for comparison using calipers.

# 5 Results

Expression of mRNA after Infection with AdCMV-p53 and AdCMV-p53-FLAG Virus. Both Tu-138 and MDA 686-LN were examined for expression of p53 mRNA. Total RNA was isolated after adenovirus infection. Northern blot analysis was performed. Similar levels of exogenous AdCMV-p53 mRNA were detected between AdCMV-p53 and AdCMV-p53-FLAG infected cells. The level of p53 mRNA expression following infection with AdCMV-p53 and AdCMV-p53-FLAG were comparable for Tu-138 and for MDA 686-LN. Variation of intensity is felt to be related to loading dose. Endogenous expression of p53 mRNA is seen in lanes 2 and 3 in the mutated p53 cell line, Tu-138. There was no significant endogenous p53 mRNA expression in the MDA 686-LN cell line, which is wild-type for the p53 gene. These data suggest that the AdCMV-p53-FLAG virus, like the AdCMV-p53 virus, is successfully transduced and efficiently transcribed. Northern analysis did not reveal evidence of AdCMV-p53 DNA contamination.

**Expression of Exogenous** *p53* **Protein in AdCMV-***p53* **and AdCMV-***p53***-FLAG Infected SCCHN Cell Lines.** Western blot analysis was performed to compare the amount of protein expressed by the AdCMV-*p53* infected and AdCMV-*p53*-FLAG infected cells. Protein bands were identified using the monospecific *p53* antibody (PAb1801) and the anti-FLAG M2 antibody (IB13025) on two simultaneously run gels. Using the *p53* antibody (pAB1801), a similar high level of *p53* protein expression in both cell lines that were infected with the AdCMV-*p53* and AdCMV-*p53*-FLAG. The Tu-138 and MDA 686-LN cells infected with the Ad-CMV-*p53*, were also tested. No change in *p53* protein expression was noted in either the replication defective adenovirus infected cells or in the mock infection group. When a similarly executed gel that was probed with the mouse anti-FLAG M2 antibody. The level of *p53*-FLAG protein expression appeared to be similar to that expressed following *p53* antibody probing, but

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no detectable band was noted in those cells infected with the AdCMV-p53 virus. The mock and DL312 infected cells exhibited no detectable level of the immunoreactive p53 or FLAG protein in either cell line.

Effect of AdCMV-p53 and AdCMV-p53-FLAG on SCCHN Cell Growth In Vitro.

The cytotoxic effect of wild-type *p53* therapy in Tu-138 and MDA 686-LN cell lines has been detailed above. The Tu-138 cell line has an endogenously mutated *p53* gene and the MDA 686-LN cell line possesses the wild-type *p53* gene. This study sought to determine if any difference in efficacy would be seen following recombination of the AdCMV-*p53* virus by inserting the FLAG sequence.

Cells infected with the replication defective adenovirus had a similar growth rate to the mock infection cells. A mild cytotoxic effect may be seen with the replication defective adenovirus (FIG. 7A). In contrast, those cells infected with either the AdCMV-p53 or AdCMV-p53-FLAG experienced virtual total tumor cell death by day three. Histologic examination revealed bleb formation by the plasma membrane which is the characteristic feature of apoptosis and has been characterized as the mechanism of cell death in AdCMV-p53 infected SCCHN cell lines (Example 1). As noted above, the effect was more prominent for the Tu-138 cell line (mutated p53) than it was for the MDA 686-LN cell line (wild-type p53). Growth curve assays were reproducible in three repeated studies without a significant difference being noted between the effect of the AdCMV-p53 and the AdCMV-p53-FLAG viruses, suggesting that the addition of the FLAG peptide did not affect the ability of p53 in suppression of cell growth.

Immunohistochemical Staining of SCCHN Cell Lines Infected with Adenovirus.

Infected cell monolayers were compared for expression of the *p53* and *p53*-FLAG protein using standard immunohistochemical techniques. Neither the *p53* nor FLAG protein could be clearly identified in the mock infection of DL312 infected cells in the MDA 686-LN cell line. However, in Tu-138, which has a mutated *p53* gene, endogenous staining for *p53* was positive. When cells were infected with the AdCMV-*p53* virus a strong staining was noted in both cell

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lines. Visual inspection of those cells infected with the AdCMV-p53-FLAG virus showed identical intensity of staining and number of positive cells with PAb1801 for antibody as compared to the cells infected with the AdCMV-p53 virus. The cells infected with the AdCMV-p53-FLAG virus also showed strong immunohistochemical positivity with the M2 FLAG antibody. The quality of staining was different though, both within the nucleus and to a lesser degree in the cytoplasm.

In Vivo Suppression of Growth. Dose response studies using  $10^8$ ,  $10^9$ , and  $10^{10}$  plaque forming units (pfu) of AdCMV-p53-FLAG virus compared to a control flap that was either PBS or DL312 were performed using the microscopic model method described in Example 1 on the Tu-138 cell line. The mean tumor size for the mock infection was  $1205 + /-205 \text{ mm}^3$ . Tumor size decreased in a linear fashion with increasing concentration of virus used in the molecular intervention. Mean tumor size was  $637 + /-113 \text{ mm}^3$ ,  $392 + /-109 \text{ mm}^3$ , and  $193 + /-74 \text{ mm}^3$  for those flaps treated with  $10^8$ ,  $10^9$ , and  $10^{10}$  pfu of the AdCMV-p53-FLAG respectively. Each animal was compared against itself using a paired t test and a significant dose response effect was noted at p<0.05 in all comparisons except between the flap treated with  $10^9$  and  $10^{10}$  pfu. Clearly, the greater the amount of virus, the greater the tumor growth inhibition visualized. In an additional study the effects of AdCMV-p53 were compared to that of the AdCMV-p53-FLAG. No significant difference in activity was noted.

Immunohistochemical Demonstration of Exogenous Tumor Suppression Effect in the Microscopic Residual Disease Animal Model. After proving comparable in vitro and in vivo AdCMV-p53-FLAG, the inventor applied of the activity the AdCMV-p53 and immunohistochemical techniques to demonstrate p53-FLAG fusion protein product in vivo. Using the Tu-138 and MDA 686-LN cell lines, microscopic residual disease flaps were harvested 48 hours after treatment, fixed in formalin, and paraffin-embedded. On neighboring sections of tumor cells treated with the AdCMV-p53-FLAG virus, staining for both the p53 and FLAG protein were applied. Staining intensity and the number of cells staining positively was directly proportional to the amount of virus used in the infection. Controls were negative for staining

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with both *p53* and FLAG antibodies in the MDA 686-LN cells. Endogenous staining for *p53* was noted in the Tu-138 tumor cells. A histologic specimen stained with Hematoxylin and Eosin, the *p53* antibody, and the FLAG antibody. The characteristic cytoplasmic staining with the FLAG M2 antibody contrasted with the intranuclear staining of the *p53* antibody. This is the first time that the FLAG M2 antibody has been proven to be effective on paraffin-embedded fixed tissue. The staining demonstrates that the tumor suppressive effect is directed by the exogenous therapy, and that in an *in vivo* model one can identify the exogenous therapy using the applied FLAG system.

In conclusion, it is clear that the co-delivery of the FLAG protein along with the desired gene therapy offers potential utility as a marker of gene therapy. The invention clearly shows that it was simultaneously promoted along with the *p53* gene and that expression of the messenger RNA and protein were not decreased. More importantly, the biologic activity of the delivered tumor suppressor gene was not altered. For the first time, the FLAG antibody was proven effective when immunohistochemical analysis was performed on formalin-fixed paraffinembedded tissue. These factors suggest the utility of this novel protein as a tracer in further gene therapy studies.

# **EXAMPLE 5**

Treatment of Squamous Cell Carcinoma of the Head and Neck Using p53-Adenovirus

# Patient A

A 53 year-old male patient presents with an inoperable SCCHN tumor of the head. The tumor mass is approximately 6.5 cm diameter. Following examination of bone marrow function, platelet count and renal function, the patient receives a first treatment with 10<sup>8</sup> infectious particles of an adenovirus-*p53* expression construct, diluted in sterile phosphate buffered saline, via eight distinct intratumoral injections (total volume 10 ml). Every three days, the patient receives an identical treatment, until a total of six treatments have been given.

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Three days after the sixth treatment, the tumor is examined and found to be > 4.0 cm in diameter. Histologic examination shows considerable cell fragmentation at the tumor margin. A second course of six treatments in undertaken, following which the tumor is found to be > 2.0 cm in diameter and necrotic. The patient continues to receive one a week treatments for three months, at which time the tumor is no longer evident.

# Patient B

A 44 year-old female patient presents with an operable SCCHN tumor of the neck. The tumor mass is approximately 2.5 cm diameter. Following examination of bone marrow function, platelet count and renal function, the patient receives a first treatment with 5 X 10<sup>7</sup> infectious particles of an adenovirus-*p53* expression construct, diluted in sterile phosphate buffered saline, via three distinct intratumoral injections (total volume 3 ml). Every three days, the patient receives an identical treatment, until a total of six treatments have been given.

Three days after the sixth treatment, the tumor is excised. The tumor bed is bathed in 6 ml of sterile phosphate buffered saline for 60 min. The inoculum is removed, the wound closed, and a drain left in the tumor bed. On days 4, 7, 10 and 14 following surgery, 5 X 10<sup>7</sup> infectious particles of an adenovirus-*p53* expression construct, diluted in sterile phosphate buffered saline (total volume 3 ml) are infused via the drain. After contacting the tumor bed for two hours, the inoculum is removed by suction. Sixth months following termination of treatment no primary, local or regional tumors are observed.

#### EXAMPLE 6

# Wild-Type P53 Gene Transfer Via An Adenoviral Vector In A Phase I Trial Of Patients With Advanced Recurrent Head And Neck Squamous Carcinoma

An adenoviral vector containing the normal "wild-type" p53 gene was delivered in logorithmically increasing doses in patients with biopsy-confirmed recurrent squamous carcinoma of the head and neck. Direct tumoral injections were performed three times weekly

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for two consecutive weeks. Patients were stratified into two groups: 1) resectable recurrent disease, 2) unresectable recurrent disease.

Those patients stratified into the resectable disease group underwent total gross surgical resection of their recurrent neoplasm 72 hours following the sixth gene transfer events over a 2-week period. The adenoviral vector also was delivered intraoperatively and 72 hours following the surgical procedure via retrograde catheter infusion. Unresectable patients underwent repeat gene transfer attempts via direct tumoral injections monthly over two week cycles until disease progression or deterioration in patient performance status was observed. The safety of this treatment was monitored by close hospital observation, biopsies to assess gene transfer efficiency, bodily fluid analysis for shed vectors and necropsy analysis.

# **Methods**

Study Subjects. Twenty-one patients with advanced recurrent squamous carcinomas of the upper aerodigestive tract with an Eastern Cooperative Oncology Group performance status 2 patients were entered into one of two study arms consisting of patients with resectable (Group 1) or non-resectable (Group 2) recurrent malignancies. The characteristics of the study subjects and dosage of adenoviral vector are shown in Tables 6 and 7, respectively. The women all had negative pregnancy tests and all patients used contraceptive methods. Informed consent was obtained from all patients prior to study entry.

Gene-Transfer Vector. The present study employed a replication-defective adenovirus serotype 5 vector with an enhancer (cytomegalovirus)-promoter designated Ad5CMVp53. Three lots of adenoviral vector with plaque-forming units (PFU) ranging from 10<sup>9</sup> to 10<sup>11</sup> were produced by good manufacturing practices at Magenta, Inc. and Introgen Therapeutics, Inc. and shipped frozen (-70°C) to the University of Texas, M.D. Anderson Cancer Center. Each lot was efficacious for transduction utilizing western blotting as well as an *in vitro* tumor cell suppression growth assays. Vector was thawed and diluted in phosphate-buffered saline (vehicle) immediately prior to gene transfer and transported to patient rooms at 4°C.

Table 6. Patient Characteristics

1         Cancer         Gancer         Therapies         Transfer         Arm           2         58         M         Laryn         Moderately         WT         surgery, xrt, scial mass         1           3         56         M         Pyriform         Poorly diff.         WT         surgery, xrt, submental         1           4         42         F         Base of sorly diff.         WT         surgery, xrt, submental         2           5         M         Pyriform         Poorly diff.         WT         surgery, xrt, submental         2           4         42         F         Base of sorly diff.         WT         surgery, xrt, sight neck         2           5         71         M         Unknown         Poorly diff.         WT         surgery, xrt, sight neck         1           6         43         M         Cervical         Poorly diff.         WA         surgery, xrt, settence         1           7         6         43         M         Cervical         Poorly diff.         WA         surgery, xrt, settence         2           8         7         6         A         chemo         mass           9         7         6         chemo<	Pt. No.	Age	Gender	Primary	Pathology	p53	Prior Failed Site of Gene	Site of Gene	*Study
31 F Floor of Moderately MAA surgery, xrt, Facial mass  Mouth diff. SCC chemo  assopharynx  diff. SCC chemo  Aiff. SCC chemo  assopharynx  differential check  Sinus differential check  Base of Poorly diff. WT surgery, xrt Right neck  tongue SCC chemo mass  SCC chemo mass  SCC chemo mass  scophasgus SCC chemo mass  diff. SCC chemo chemo chemo mass  diff. SCC chemo Left neck  diff. SCC chemo Left neck  scophasgus SCC chemo mass  diff. SCC chemo Left neck  mass  diff. SCC chemo Left neck  mass  diff. SCC chemo Left neck  mass				Cancer		genotype	Therapies	Transfer	Arm
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58         M         Larynx         Moderately         WT         surgery, xrt         Submental mass, adiff. SCC           56         M         Pyriform         Poorly diff.         MAA         surgery, xrt         Left neck           42         F         Base of tony diff.         Poorly diff.         WT         surgery, xrt         Right neck           71         M         Unknown         Poorly diff.         WT         surgery, xrt         Left neck           43         M         Cervical,         Poorly diff.         MAA         surgery, xrt,         Peristomal           43         M         Cervical,         Poorly diff.         MAA         surgery, xrt,         Peristomal           43         M         Cervical,         Poorly diff.         MAA         surgery, xrt,         Peristomal           43         M         Cervical,         Poorly diff.         MAA         surgery, xrt,         Peristomal           43         M         Cervical,         Poorly diff.         MAA         surgery, xrt,         Peristomal           43         M         tonsil         Moderately         WT         xrt, chemo         Left neck				Mouth	diff. SCC		chemo		
56         M         Pyriform         Poorly differential         MAA         surgery, xrt         Left neck           42         F         Base of tongue         Poorly diff.         WT         surgery, xrt         Right neck           71         M         Unknown         Poorly diff.         WT         surgery, xrt         Left neck           85         M         Unknown         Poorly diff.         WT         surgery, xrt         Left neck           843         M         Cervical, corright         Poorly diff.         MAA         surgery, xrt         Peristomal           943         M         Cervical, corright         Poorly diff.         MAA         surgery, xrt         Peristomal           943         M         Cervical, corright         Poorly diff.         MAA         surgery, xrt         Peristomal           943         M         tonsil         Moderately         WT         strt, chemo         Left neck           943         M         tonsil         Moderately         WT         strt, chemo         Left neck	7	58	M	Larynx	Moderately	WT	surgery, xrt	Submental	н
56 M Pyriform Poorly MAA surgery, xrf Left neck Sinus differential mass 42 F Base of Poorly diff. WT surgery, xrf Right neck tongue SCC chemo mass  71 M Unknown Poorly diff. WT surgery, xrf Left neck SCC chemo mass SCC chemo mass SCC chemo mass chemo mass SCC chemo least chemo mass SCC chemo least chemo least chemo mass SCC chemo least chemo least chemo least neck SCC chemo least chemo least chemo least neck SCC chemo least chemo least chemo least neck SCC chemo least chemo least neck SCC chemo least neck SC					diff. SCC			mass,	
56MPyriformPoorlyMAAsurgery, xrtLeft neck42FBase of tonly diff.WTsurgery, xrtRight neck71MUnknownPoorly diff.WTsurgery, xrtLeft neck71MUnknownPoorly diff.WTsurgery, xrtLeft neck43MCervical,Poorly diff.MAAsurgery, xrtPeristomal63McevphasgusSCCchemomass63MtonsilModeratelyWTxrt, chemoLeft neck								nasopharynx	
Hander Sinus differential more boorly diff. WT surgery, xrt Right neck tongue SCC mass mass and tongue SCC chemo mass diff. SCC chemo mass mass sCC chemo ma	m	56	M	Pyriform	Poorly	MAA	surgery, xrt	Left neck	2
42 F Base of Poorly diff. WT surgery, xrt Right neck tongue SCC mass  71 M Unknown Poorly diff. WT surgery, xrt, Left neck SCC chemo mass  8 SCC chemo mass  9 SCC chemo mass  9 SCC chemo mass  1 SCC chemo mass  1 SCC chemo mass  1 SCC chemo mass  2 SCC chemo mass  3 SCC chemo left neck chemo mass  4 SCC chemo left neck mass  4 SCC chemo left neck mass  4 SCC chemo left neck mass  6 SCC chemo left neck mass  7 SCC chemo left neck mass  8 SCC chemo left neck mass				Sinus	differential			mass	
M Unknown Poorly diff. WT surgery, xrt, Left neck SCC chemo mass M Cervical, Poorly diff. MAA surgery, xrt, Peristomal esophasgus SCC chemo mass M tonsil Moderately WT xrt, chemo Left neck mass	4	42	ц	Base of	Poorly diff.	WT	surgery, xrt	Right neck	2
M Unknown Poorly diff. WT surgery, xrt, Left neck SCC chemo mass M Cervical, Poorly diff. MAA surgery, xrt, Peristomal scophasgus SCC chemo mass M tonsil Moderately WT xrt, chemo Left neck diff. SCC mass				tongue	SCC			mass	
SCC chemo mass M Cervical, Poorly diff. MAA surgery, xrt, Peristomal esophasgus SCC chemo mass M tonsil Moderately WT xrt, chemo Left neck diff. SCC mass	\$	71	M	Unknown	Poorly diff.	WT	surgery, xrt,	Left neck	1
M Cervical, Poorly diff. MAA surgery, xrt, Peristomal esophasgus SCC chemo mass M tonsil Moderately WT xrt, chemo Left neck diff. SCC mass					SCC		chemo	mass	
esophasgus SCC chemo mass M tonsil Moderately WT xrt, chemo Left neck diff. SCC mass	9	43	M	Cervical,	Poorly diff.	MAA	surgery, xrt,	Peristomal	7
M tonsil Moderately WT xrt, chemo Left neck diff. SCC mass				esophasgus	SCC		chemo	mass	
	7	63	M	tonsil	Moderately	WT	xrt, chemo	Left neck	2
					diff. SCC			mass	

Table 6 continued:

-	2	1	-	8	2	7
Base of tongue	Peristomal Recurrence	Hypopharyng eal mass	Base of tongue	Left retromolar	trigone Posterior ora tongue &	tongue base Floor of mouth
surgery, xrt	surgery, xrt	surgery, xrt	xrt, chemo	surgery, xrt	surgery, xrt	surgery, xrt
MAA	WTT	MAA	MAA	WT	WT	WT
Moderately diff. SCC	Well diff.	Moderately diff. SCC	Poorly diff. SCC	Moderately diff. SCC	Moderately diff. SCC	Poorly diff. SCC
Base of tongue	Larynx	Larynx	Base of tongue	Floor of mouth	Base of tongue	Floor of mouth
M	M	M	M	ш	M	ίτι
46	57	58	46	57	99	84
∞	6	10	11	12	13	41

	7		-		2			2			П		2		2	
	Facial mass		Supraclavicul	ar mass	Base of	tongue,	tonsillar foss	Facial neck	mass		Submental	mass	Neck mass		Neck Mass	
	surgery, xrt,	chemo	xrt		surgery, xrt,	chemo		chemo			surgery, xrt,	chemo	surgery, xrt,	chemo	chemo, xrt	
	WT		MAA		WT			WT			WT		WT		WT	
	Moderately	diff. SCC	Poorly diff.	SCC	Moderately	diff. SCC		Moderately	diff. SCC		Poorly diff.	SCC	Poorly diff.	SCC	Well diff.	SCC
	Mandible	alveolar ridg	Larynx		Larynx			L. Lateral	pharyngeal	wall	Unknown		Tongue		Base of	tongue
	M		Щ		M			M			M		M		M	
continued:	63		92		99			56			53		55		99	
Table 6	15		16		17			18			19		20		21	

Table 7. Vector Dosing and Presence or Absence of Adenoviral Vector Nucleotides in Serum or Urine During

Treatment

Patient	PFU/Transfer	Volume	Gene Transfer Cycles	Site of Injections
		Injected		
1	10 <sup>6</sup>	6сс	2	Facial mass
2	$10^6$	3cc	2	Submental mass,
				neopharynx
3	$10^{6}$	3cc, 4cc	1	Left neck mass
4	$10^{6}$	3cc, 5cc	1 + 3	Right neck mass
5	$10^{6}$	1.5cc	1	Left neck mass
6	$10^6$	3cc	1	Peristomal mass
7	10 <sup>7</sup>	2cc	4 + 2	Left neck mass
8	$10^7$	3cc	1	Base of tongue
				mass
9	$10^7$	1.5cc	1	Peristomal
				recurrence
10	$10^{8}$	1.5cc	1	Hypo-pharyngx
				mass
11	$10^{8}$	3cc	1	Base of tongue
12	108	1.5cc	3	Left retromolar
				trigone
13	10 <sup>9</sup>	1.5cc	6	R. Posterior
10				tongue & tongue
				base
14	$10^{9}$	1.5cc	1 + 1	Floor of mouth
15	109	3cc	1	Left facial mass
16	10 <sup>9</sup>	1.5cc	1	L. Supraclavicular
10	10	1.500	-	mass
17	$10^{9}$	1.5cc	1	Base of tongue,
17	10	1.500	_	tonsillar fossa
18	10 <sup>9</sup>	3cc	4	Left facial mass
19	$10^{9.5}$	1.5cc	1	Submental
17			•	mass
20	$10^{9.5}$	3cc	3	Left neck mass
21	$10^{9.5}$	1.5cc	3	Right neck mass

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**Dosage**. The adenoviral vector was administered to five cohorts of patients each, and logarithmically increasing doses. Dosage increases were established following two weeks of observation of the last patient treated at the prior dose. Following entry of the first six patients into the study, three patients were entered at each dosage level independent of the resectable or unresectable group the patient had been stratified to. The dose of biologic vector is described in terms of the total dose (in plaque-forming units). The estimated number of vectors administered per malignant epithelial cell was not approximated. The total volume of administration is shown in Table 7. The volume of adenoviral vector injected into the solid malignancies was determined by the clinical and radiographic estimated tumor volume. The vector was directly injected into the recurrent squamous carcinomas under direct vision and by manual palpation. Injections were spaced at 1 centimeter increments across masses. After gene transfer, the subject remained under close observation for at least 1-1/2 hours. Respiratory and body secretion isolation were maintained for 72 hours following the final gene transfer of vector.

**Detection of Vector**. The urine and serum samples were monitored for disseminated adenoviral vector utilizing viral culture of 293 cells as well as polymerase chain reactions (PCR) using primers that amplify the E1b region of the adenovirus and the 5' end of the wild-type p53 gene which were specific to the vector. PCR products were then southern transferred to improve viral detection to 1-5 viral particles as well as verify PCR product specificity. Positive and negative controls were assayed in each reaction.

Safety. The symptoms, vital signs, blood counts were monitored, and the patients were physically examined and photographically documented daily. Chest radiography, blood chemistry testing, and performance status analysis were performed at the beginning of each treatment cycle. Serum titers of adenoviral antibody were measured before and following each cycle of gene transfer. Three days following the sixth gene transfer of the first cycle, tumor biopsies (or surgical resection) were obtained. Specimens were stored as snap-frozened, pathologically embedded specimens, as well as formalin-fixed specimens in every instance.

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Extraction of Nucleic Acids from Serum or Urine. Ad5CMV-p53 adenoviral DNA was extracted from 0.5-ml aliquots of serum or urine by a method modified from Cunningham *et al.*, (1995). Briefly, distilled water was added to make 1 ml, and they were precipitated with 30% polyethyleneglycol (PEG). Since SDS alone was not sufficient to release the viral DNA from its particle, proteinase K was added to the SDS at 50°C for 2-16 hours following the PEG precipitation (Norder *et al.*, 1990). The samples were extracted with phenol, and viral DNA was precipitated with ethanol in the presence of glycogen (Cunningham *et al.*, 1995). The precipitated DNA was recovered by centrifuging at 14000 g for 10 minutes at 4°C, resuspended in 0.3 ml of distilled water, and reprecipitated with ethanol. The DNA pellet was rinsed with 70% ethanol, vacuum-dried, and dissolved in 10 μl of distilled water. Samples were either analyzed immediately or stored at -20 until used. Extraction of nucleic acids was performed under biologic safety cabinet hoods to prevent possible cross-contamination of specimens.

PCR Reactions on DNA Isolated from Serum Samples. Primers were designed for specific amplification of the p53 gene from the adenoviral vector. The upper primer (5'-CACTGCCCAACAACACCA-3', SEQ ID NO:5) corresponds to the 3' end of the p53 gene and the lower primer (5'-GCCACGCCCACACATTT-3' SEQ ID NO:6) corresponds to the E1B region of adenovirus type 5 (nucleotides 3517 to 3533 of the wild-type sequence). Each PCR reaction tube contained 0.2 mM of each oligonucleotide, 0.4 mM dNTPs, 1X TaqPlus Long low salt buffer (from Stratagene), 0.6 µl TaqPlus Long (5U/ml) (from Stratagene), and 5 µl of test The samples were placed in an MJ Research Peltier Thermal Cycler (PTC-200) DNA. programmed for 93°C for 3 minutes, with the following three-step profile: 93°C for 30 seconds, 65°C for 45 seconds and 72°C for 45 seconds for a total of 30 or 35 cycles. 5 ul of 6X loading buffer (0.25% bromophenol blue, 0.25% xylene cyanol FF and 15% Ficoll (Type 400; Pharmacia) in water) was added to each tube at the end of the PCR run and loaded onto a 1% agarose, 1X TBE gel containing ethidium bromide (.6 µg/ml). The samples were electrophoresed at 100V for 1-1.5 hours and then photographed under UV light.

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Polymerase Chain Reaction (PCR). Only 5 µl of prepared DNA could be used in a single polymerase chain reaction. For serum, the PCR was performed in a 20-µl volume containing 2 mM MgCl<sub>2</sub>, 50 mM KCl, 0.1% Triton X-100, 200 µM each of deoxyribonucleoside triphosphates (dNTP), 10 mM Tris-HCl (pH 9.0), 5 µM each of the primers, and 1.7 units of Tag DNA polymerase (Promega). The reactions were carried out at 94°C for 30 sec, 58°C for 30 sec, and 72°C for 60 sec for 35 cycles, followed by a 10-min extension at 72°C. The PCR primers were selected from the sequence of the Ad5CMV-p53 with the sense primer located at the 3' end of the p53 cDNA (5'-GCCTGTCCTGGGAGAGACCG-3', SEQ ID NO:7), and the antisense (5'region adenovirus of type primer was selected from the E1B CCCTTAAGCCACGCCCACAC-3', SEQ ID NO:8). The PCR product (an 838-bp fragment) was separated on 1% agarose gel. The same PCR product was subcloned into pCR-Script vector (Stratagene), sequenced, and the gel purified insert was used as a probe to detect the PCR product. For urine, the PCR was performed in a 20-µl volume containing 2 mM MgSO<sub>4</sub>, 10 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 10 mM KCl, 0.1% Triton X-100, 20 mM Tris-HCl (pH 8.8), 0.1 mg/ml bovine serum albumin, 200 µM each of deoxyribonucleoside triphosphates (dNTP), 5 µM each of the primers, and 2.5 units of TaqPlus long DNA polymerase (Stratagene). The reactions were carried out at 93°C for 60 sec, and then 93°C for 30 sec, 65°C for 45 sec, and 72°C for 45 sec for 35 cycles, followed by a 10-min extension at 72°C. The PCR primers were selected from the sequence of the Ad5CMV-p53 with the sense primer located at the 3' end of the p53 cDNA (5'-CACTGCCCAACAACACCA-3', SEQ ID NO:9), and the antisense primer was selected from the E1B region of adenovirus type 5 (5'-GCCACGCCCACACATTT-3', SEQ ID NO:10). The PCR product (an 724-bp fragment) was separated on 1% agarose gel.

**Southern Blot Analysis.** In the Southern blot used to verify the PCR product's specificity, the DNA in the gel was denatured and neutralized before blotting to nylon membrane (Hybond-N+, Amersham) by capillary absorption. The membrane was prehybridized for 15 min at 65°C in Rapid-hyb buffer (Amersham) and hybridized in the same buffer containing <sup>32</sup>P-

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labeled probe for 1-2 h. The membrane was washed in 0.1 x SSC and 0.1% SDS at room temperature twice, and again twice at 65°C (15 min per wash). The washed membrane was exposed to x-ray film at -70°C for 1-16 h with intensifying screen.

Controls and scoring of test samples. The following controls were included with every batch of samples. At the DNA isolation step, two "negative" serum controls (pooled and aliquoted serum from Introgen employees), and two positive controls consisting of negative serum spiked with 10pfu or 100 pfu of AdCMV-p53 virus were employed. This was done to obtain a window of sensitivity where the 10 (and 100) pfu controls were positive, but the negative controls were negative. If the negative controls were positive, the PCR was repeated with only 30 cycles. If the 10 pfu control was negative, the DNA was further purified with an additional ethanol precipitation, and the PCR was repeated. The above two steps always placed the experimental parameters into the appropriate sensitivity window.

At the PCR stage, a positive control of 1 ng of AdCMV-p53 DNA (isolated from a clinical lot), and a negative (H<sub>2</sub>0) control were employed. The PCR of the batch was repeated if any of the controls were false. There were no failed negative PCR controls.

For confirmation of putative positives, the DNA was re-isolated from serum (along with several temporally adjacent samples), and the PCR of this DNA repeated. Samples were scored as positive only if the results could be reproduced. Those samples which were positive in one of two sample analyses were considered negative for reporting purposes. Putative positives that could not be repeated (due to lack of further unprocessed specimen) were omitted from the database.

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Measuring Efficacy of Gene Transfer. Surgically removed tissue specimens were placed in cryovials, immediately flash frozen, and then stored in liquid nitrogen storage until use. Frozen samples were decanted into the orifice of a stainless steel Bessman tissue pulverizer

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(Spectrum, Houston, TX) that had been pre-cooled by immersion in liquid nitrogen. Tissue was crushed to a fine powder by striking the Bessman pestle with a steel hammer five to ten times. Pulverized tissue was transferred to a glass tissue homogenizer (Fisher Scientific, Pittsburgh, PA) containing 1 ml TRI reagent (Molecular Research, Cincinnati, OH) per 50 mg tissue, and homogenized by five to ten up and down strokes with a Teflon pestle.

Following homogenization, RNA was isolated according to the instructions provided with the TRI reagent. Briefly, homogenates were transferred to polypropylene centrifuge tubes (Molecular Research) and stored for 5 min at room temperature prior to the addition of chloroform (0.2 ml per 1 ml of TRI reagent). Samples were then mixed vigorously, incubated an additional 15 min at room temperature, and centrifuged at 12,000 x g for 15 min at 4°C to separate the RNA containing aqueous layer from the phenol-chloroform phase. Isopropanol was added to the aqueous phase and RNA was precipitated by incubating at room temperature for 15 min. RNA pellets were recovered by centrifugation at 12,000 x g for 15 min at 4°C, washed once with 75% ethanol, air dried, dissolved in diethyl pyrocarbonate (DEPC)-treated water and quantitated by measuring the absorbance at 260 nm. Contaminating DNA was removed by incubating up to 50 µg RNA with 60 U DNase I (Pharmacia, Piscataway, NJ) for 25 min at 37°C in a total reaction volume of 260 ul. RNA was then extracted with phenol:chloroform, ethanol precipitated, washed once with 75% ethanol, pelleted by centrifugation at maximum speed in a microfuge for 15 min at 4°C, air dried, resuspended in DEPC-water, and stored at -80°C. The quality of RNA was assessed by running samples on an ordinary non-denaturing 0.8% agarose gel and visualizing the 28S and 18S ribosomal bands by ethidium bromide staining. eliminate cross-contamination between samples and to minimize RNase activity, all re-usable instruments employed for RNA isolation were soaked in a 2% Liqui-Nox (Fisher Scientific) detergent solution for a minimum of 5 minutes, scrubbed free of debris, transferred to a 10% bleach solution for 3 minutes, rinsed thoroughly with deionized water, sprayed with 100% ethanol, dried, immersed in chloroform, and dried again before use.

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Reverse transcription was performed using 1.5 µg total cellular RNA in a 23.5 µl reaction mix containing 111 ng random hexamers (Gibco BRL, Grand Island, NY), 40 units RNase inhibitor (Boehringer Mannheim, Indianapolis, IN), 0.4 mM of each dNTP (Perkin Elmer, Foster City, CA), and 300 units of Superscript II RNase H reverse transcriptase (Gibco BRL) in 1 x RT buffer (50 mM Tris pH 8.3, 75 mM potassium chloride, 3 mM magnesium chloride, and 20 mM dithiothreitol). RNA and random hexamers were heated to 70°C for 10 min and cooled on ice before the rest of the reaction mix was added. The reaction was incubated at 25°C for 5 min with 200 units reverse transcriptase, and then an additional 10 min at 25°C following addition of another 100 units reverse transcriptase, to facilitate primer annealing, before incubating at 42°C for 50 min. The RT-reactions were terminated by heat inactivating the reverse transcriptase for 15 min at 70°C. RNA complementary to cDNA was removed by digestion with 1 unit RNase H (Boehringer Mannheim) for 20 min at 37°C. RNA from the head and neck squamous cell carcinoma (HNSCC) line TU167 infected with recombinant adenovirus Ad5CMV-p53 (multiplicity of infection of 100:1) was used as a positive control for detecting virally transcribed p53, and TU167 cells infected with the variant adenovirus vector dl312 (1) that does not contain the p53 transcriptional unit was used as a negative control.

To detect the Ad5CMV-*p53* transcript, PCR was performed in a reaction volume of 30 μl containing 0.2 mM of each dNTP, 1.5 mM magnesium chloride, 1 unit taq polymerase (Promega, Madison, WI), and 0.5 mM of each primer CMV2 (5' GGTGCATTGGAACGCGGATT, SEQ ID NO: 11) and P53EX3 (5' GGGGACAGAACGTTGTTTTC, SEQ ID NO:12) in 1 x PCR buffer (50 mM potassium chloride, 10 mM Tris pH 9.0, 0.1% Triton X-100). Primers CMV2 and P53EX3 amplify a 295-base fragment specific to the adenovirus derived p53 transcript. PCR conditions for detecting Ad5CMV-*p53* transcripts were as follows:

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94°C for 1 min, followed by 94°C for 30 sec, 58°C for 40 sec, 70°C for 1 min for 35 cycles, and 70°C for 10 min extension time.

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To ensure that the product amplified during PCR was detecting mRNA and not contaminating DNA in the RNA preparation, PCR was also performed using RT products from parallel reactions in which no reverse transcriptase was added.

RT-PCR specific for glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was performed in order to check the integrity of the RT reactions. A 3 µl volume of the RT reaction was diluted in 30 µl of PCR mix containing 0.2 mM of each DNTP, 2 mM magnesium chloride, GAPDH1 unit polymerase, and 0.5 mMof each primer taq NO:13) GAPDH2 ACGGATTTGGTCGTATTGGG, **SEQ** ID and (5' TGATTTTGGAGGGATCTCGC, SEQ ID NO:14) in 1 x PCR buffer. The GAPDH primers span 3 exons in the human GAPDH gene and amplify a 231-base product specific for mRNA. PCR conditions to detect GAPDH were as follows:

94°C for 1 min, followed by 94°C for 30 sec, 60°C for 12 sec, 72°C for 1 min for 35 cycles, and 72°C for 7 min extension time.

PCR was performed using a Perkin Elmer Gene Amp 9600 thermocylcer, and all primers were commercially synthesized (Genosys, Woodlands, TX).

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Immunohistochemical Determination of Intratumoral Gene. Immunoperoxidase studies were performed on formalin-fixed, paraffin-embedded tissue sections using the avidin-biotin-peroxidase complex (ABC) method (1). The specimens were cut 3-4µm thick, deparaffinized in xylene, and rehydrated in descending grades (100-70%) of ethanol. Endogenous peroxidase activity was blocked with 3% hydrogen peroxide in methanol. After several washes in distilled water and phosphate buffered saline (PBS), section were incubated with a 1:10 dilution of normal horse serum to minimize background staining. This was followed by overnight incubation at 4°C with monoclonal antibodies to p53 (DO-1, Oncogene Science, Inc., Uniondale, NY; 1:80 dilution) and p21 (Oncogene Science, Inc., 1:100). Peroxidase staining procedure was done using ABC Elite kits (Vector Laboratories, Burlingame, CA). The immunostaining reaction was visualized using 0:05% 3.3'-diaminobenzidene in Tris-HC1 buffer containing 0.01% hydrogen peroxide, pH 7.6. Sections were counterstained with 0.01% toluidine blue and mounted in permount. Scoring was performed by counting positive nuclear staining in 200 cells of 10 consecutive high power fields by two independent observes.

TUNEL Assay for DNA Fragmentation. The TUNEL assay was performed using the Apoptag<sup>TM</sup> PLUS kit (Oncor, Gaithersburg, M.D.) according to instructions provided by the manufacturer. Slides were counterstained with .4% methelene green. Corresponding hematoxilin and eosin stained slides were evaluated for the presence of inflammatory cell infiltrates and were graded on a scale of 1-4.

Cytopathic Effect Assay Procedure. Patient urine samples also were monitored for the presence of Ad5p53 by an assay in which any virus in the sample is allowed to infect a recipient cell monolayer. Those cells are monitored for the appearance of a cytopathic effect (CPE): the cells round up and detach from the surface. The patient urine samples assayed for CPE were from the first morning urines collected during the second week of the first treatment course, and on Day 0 or 1 (pretreatment). Samples of approximately 15 mls each were stored in sterile 15-ml conical tubes at -80°C until use. The IT293 cells, which form the recipient cell monolayer in this assay, are maintained in DMEM plus 10% FBS, in a 37°C humidified 10% CO<sub>2</sub> incubator. Two days prior to patient sample testing, the cells were plated at 2 x 10<sup>5</sup> per well in 12-well plates.

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At time of assay, the urine samples were thawed in an ice bath, and an aliquot was mixed 1:1 with DMEM and sterile-filtered using a 0.22 µm syringe filter. A 350 µl aliquot of this 1:1 mix was slowly added to each well after removing the growth medium. The plate was gently rocked 15 minutes later. After 30 minutes, 2.0 mis of DMEM plus 10% FBS was added to each well to dilute the sample. On day 3 (72 hours later) and day 6 of the assay, 0.5 ml aliquots of fresh medium were added to each well, to help maintain the cell monolayer for the maximum 6-7 days.

Patients samples were assayed in triplicate. The diluted pretreatment sample was assayed as is, and also spiked with 10<sup>5</sup> pfu of Ad5p53 per well to detect any urine component that might interfere with the detection of virus by this procedure. Control wells were inoculated with DMEM alone spiked with 10<sup>5</sup>, 10<sup>4</sup>, 10<sup>3</sup>, 10<sup>2</sup>, or 10<sup>1</sup> pfu per well, each in duplicate. The 10<sup>5</sup> pfu spike causes CPE under these conditions on day 2 of the assay; on each successive day, the next spiked control will show CPE. Therefore, the time at which CPE is detected in each patient sample indicates the level of Ad5p53 in that sample.

**Recombinant Competent Adenovirus.** The adenoviral p53 used in the clinical trial was tested for the presence of RCA using A549 cells by the Biotechnology Services Division of Microbiological Associates, Inc., (Rockville, Maryland).

Statistical Analysis. A single-arm study design was used. To prevent enrolling more patients than necessary in a trial of excessive toxicity was found, a Bayesian early stopping rule was implemented. The WILCOXON signed rank test and assigned test were used for comparisons between before and after treatment of the percentage of cells showing TUNEL and immunohistochemical staining, respectively. Statistical analysis was performed using the Survpac SPSS-Statistical package.

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Response and Toxicity. Survival and response were assessed in this interim analysis, but were not considered aims of this analysis. The purpose of this interim analysis was to determine transduction potential of this gene transfer strategy. Patients were evaluable for response and toxicity following a 30-day observation after one cycle of gene transfer. Toxic effects of therapy were evaluated according to the National Cancer Institute's common toxicity criteria. (Xref.) Response to therapy was assessed by CT scan or ultrasound of the neck before each course of treatment. Patients were evaluable for response if they received at least one course of therapy followed by an appropriate documentation of response. Those patients undergoing surgical resection of their recurrent tumors could not be evaluated for response since surgery was performance prior to a 30 day observation period. All CT scans were evaluated by one radiologist and ultrasounds by another. Partial response was defined as a 50 percent or greater reduction in the sum of the products of the diameters of the measureable tumor; a minor response was defined as a 25 percent to less than 50 percent reduction in the sum of the products of the diameters of the measureable lesion. Progression of disease was defined as a 25 percent or greater increase in the sum of the products of the diameters.

Survival duration was measured from time of protocol entry until expiration. Each patient's response was reviewed by a data management committee consisting of a Head and Neck Surgical Oncologist, Radiologist, and Medical oncologist.

# **Results**

**Detection of Vector.** Adenoviral-vector DNA was detected by PCR in serum as well as urine samples from patients up to 48 hours following gene transfer. The detection limit for vector is 1-5 viral particles. Viral DNA was isolated in urine among patients undergoing gene transfer at each viral dosage, but not detected after 48 hours following gene transfer. Serum detection of viral DNA increased with increasing viral dosage above 10<sup>7</sup> PFU, but was also not detectable 48 hours following gene transfer.

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Urine samples were first analyzed for infectious virus, measured CPE on cell monolayers, prior to PCR nucleotide analysis. The virus present in the urine was applied to 293 cells as described above, in order to monitor the CPE, which is observed as the cells round up and become from the petri dish. CPE was identified rarely in specimens. CPE was found late in overlay cultures (greater than six days) and these results were considered equivocable. In no instance was CPE confirmed by PCR and Southern blot transfer of the same urine sample for detected adenoviral nucleotides.

Viral DNA Analysis of other Organ Systems. PCR analysis demonstrated viral DNA greater than two months following 10<sup>9</sup> Ad5CMVp53 gene transfer in skin, cardiac muscle, lung, and testicular tissues in snap frozen autopsy specimens carefully sampled to prevent cross-contamination between tissues. Renal parenchyma, adrenal glands and pancreatic tissues did not show viral DNA sequences specific for this vector. Immunohistochemical analysis of these autopsy specimens (did not reveal evidence of wild-type p53 protein product overexpression).

Assessment of Gene Transfer. All analyses were performed after at least 1 hour of exposure of tissues to the vector. The mRNA product was detected at 4 hours and 48 hours via RT-PCR. In contrast, biopsy specimens frozen after 1 hour of exposure or less, showed no p53 mRNA. In addition, negative control samples obtained from an operative site that had not been subjected to gene transfer were also negative for the PCR product. Non-transduced and transduced biopsy specimens from four patients were analyzed by RT-PCR for the presence of Ad5CMVp53 transcribed mRNA. Transduced specimens from two patients were positive on EtBr stained gels, while no Ad5CMVp53 product was detected in any of the non-transduced specimens despite the fact that GAPDH could be amplified from all specimens. The specificity of the 295 bp PCR product from the two positive patients was confirmed by Southern Blotting. Because PCR product was not observed when reverse transcriptase was ommitted from the RT reactions, the 295 bp product detected in the figure had to have been generated from mRNA, rather than contaminating DNA.

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Immunohistochemical Analysis. All patients pre- and post-gene transfer specimens were simultaneously analyzed with positive and negative controls in each experiment. Multiple sections of each specimen were analyzed and compared with hematoxylin and eosin-stained as well as TDT end-labeled specimens. Post-vector injected biopsy specimens confirmed gene transfer in three patients whose pre-transfer biopsy specimens did not show endogenously overexpressed p53 protein. In 5 of 21 patients (27%), p21 (CIP/WAF1) was not significantly endogenously expressed; this proved informative for gene transfer in the patient post-gene transfer biopsy specimens. p53 nuclear protein overexpression was also seen in tumor-associated lymphocytes as well as stromal tumor cells, thus indicating nontumor cell transduction as well.

Serologic Antibody Response. Adenovirus serotype 5 antibodies were induced in all patients following repeat injections of the vector. However, transduction efficiency was not found to be significiantly altered comparing the initial cycle of gene transfer and subsequent cycles. Mild tumor injection site erythema was recognized beginning at 3 x 10<sup>9</sup> PFU., however these local reactions did not limit vector tolerance. No evidence of systemic hypersensitivity was found in any patients despite repeat viral dosing for as much as six consecutive months in a 10<sup>9</sup> PFU-treated patient.

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Pathologic Observations. Needle tracks have been identified in the majority of biopsy specimens, and gene product expression was demonstrated in deeper tissues beyond the injection sites. Similarly, the finding of end-labeled cells in transduced areas suggested the induction of apoptosis in tumor cells but the absence of apoptosis in stromal or inflammatory cells. Inflammatory cells were noted in patients and became prominent histologic findings among the patients who received doses of 10<sup>9</sup> PFU or greater. Interestingly, patient number

7, whose sampled expressed endogenous wild-type p53, demonstrated hemorrhagic necrosis with no evidence of viable tumor in serially sectioned surgical specimens in his recurrent 2 cm neck mass after one-cycle of gene transfer. Pathologic findings of necrosis as well as apoptosis induction were frequent observations in specimens.

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Clinical Observations. Patients have tolerated direct tumoral injections of vector with the most frequent adverse event being injection site discomfort. Evidence of local injection site erythema has been noted at 10<sup>9</sup> PFU of vector and higher although systemic evidence of hypersensitivity has not been seen despite evidence of systemic antibody titer elevations. Patients number 5, 10, and 16 remain without evidence of disease with a median follow-up of seven months. Patients 7 and 13 exhibited stable disease for 3 and 5 months, respectively in their indicator lesion area. Patient number 20 exhibited a partial response as verified by CAT Scan and ultrasound, but developed spinal and pleural metastasis requiring removal from study due to disease progression.

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In conclusion, patients with recurrent squamous carcinomas of the head and neck, adenovirally mediated gene transfer of wild-type *p53* is safe and can effectively transduce cancer and normal cells via direct tumoral injections or intraoperative instillation of vector. None of the patients exhibited toxicity that might limit vector administration or dose escalation. Transgene product expression was unaltered despite development of a systemic antibody response. The local inflammatory responses found pathologically within resected tumor specimens may, in fact, be beneficial.

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details supplementary to those set forth herein, are specifically incorporated herein by reference:

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# **SEQUENCE LISTING**

	(1) GEN	ERAL INFORMATION:
5	(i)	APPLICANT: Clayman, Gary L.
	(ii)	TITLE OF INVENTION: Methods and Compositions for the Diagnosis and Treatment of Cancer
10	(iii)	NUMBER OF SEQUENCES: 14
15	(iv)	CORRESPONDENCE ADDRESS:  (A) ADDRESSEE: Arnold, White and Durkee  (B) STREET: P.O. Box 4433  (C) CITY: Houston  (D) STATE: TX
		(E) COUNTRY: USA (F) ZIP: 77210-4433
20	(v)	COMPUTER READABLE FORM:
		(A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
25		(D) SOFTWARE: PatentIn Release #1.0, Version #1.30
	(vi)	CURRENT APPLICATION DATA:  (A) APPLICATION NUMBER: UNKNOWN  (B) FILING DATE: CONCURRENTLY HEREWITH  (C) CLASSIFICATION: UNKNOWN
30		(C) CHASSIFICATION: UNKNOWN
	(viii)	ATTORNEY/AGENT INFORMATION:  (A) NAME: Highlander, Steven L.  (B) REGISTRATION NUMBER: 37,642  (C) REFERENCE/DOCKET NUMBER: INGN:022
35		(c) RELEADINGS, BOOKET HOLDER! THE TOTAL
	(ix)	TELECOMMUNICATION INFORMATION: (A) TELEPHONE: (512) 418-3000 (B) TELEFAX: (512) 474-7577
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	(2) INFO	RMATION FOR SEQ ID NO:1:
	(i)	SEQUENCE CHARACTERISTICS:
4.5		(A) LENGTH: 2066 base pairs
45		(B) TYPE: nucleic acid
		<ul><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>
		(D) IOPOHOGI: IIIIEGI

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# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

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	CTGACAACC	T CI	rtggi	GAAC	CTI	'AGA'	CCT	AAAA	AGGAA	AT (	GTCAC	CCCF	AT CO	CCAC	ACCCI		1560
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3	TCCAATTTC	T T	CTTTC	CTTTT	TTT	TTT	TTT	TTTC	CTTTI	TC :	rttg <i>i</i>	GACT	rg go	TCT	CTTTC	3	1680
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	ATTTTCACC	C C	CCCI	TCCC	CTC	CTTCT	rccc	TTTI	TAT	ATC (	CCATT	TTT	AT AT	rcga:	CTCT	ŗ	2040
20	TATTTTACA	A TA	AAAA	CTTTG	CTC	CCA											2066
25	(2) INFOR	SEQU (A)	JENCI	FOR S E CHA NGTH: PE: a	RACT 293	reris	STICS ino a		3								
30	(xi)	(D)	TOP	RANDE POLOG E DES	Y: ]	linea	ar		ONO:	:2:							
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	Gly	Thr	Ala	Lys 20	Ser	Val	Thr	Cys	Thr 25	Tyr	Ser	Pro	Ala	Leu 30	Asn	Lys	
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					115					120					125				
	5		Cys	Thr 130	Thr	Ile	His	Tyr	Asn 135	Tyr	Met	Cys	Asn	Ser 140	Ser	Cys	Met	Gly	
	10		Gly 145	Met	Asn	Arg	Arg	Pro 150	Ile	Leu	Thr	Ile	Ile 155	Thr	Leu	Glu	Asp	Ser 160	
	10		Ser	Gly	Asn	Leu	Leu 165	Gly	Arg	Asn	Ser	Phe 170	Glu	Val	Arg	Val	Cys 175	Ala	
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¥ mg	20		Pro	Asn 210	Asn	Thr	Ser	Ser	Ser 215	Pro	Gln	Pro	Lys	Lys 220	Lys	Pro	Leu	Asp	
Line Ho Ho spine the time the time that the time the time that	25		Gly 225	Glu	Tyr	Phe	Thr	Leu 230	Gln	Ile	Arg	Gly	Arg 235	Glu	Arg	Phe	Glu	Met 240	
			Phe	Arg	Glu	Leu	Asn 245	Glu	Ala	Leu	Glu	Leu 250	Lys	Asp	Ala	Gln	Ala 255	Gly	
	30		Lys	Glu	Pro	Gly 260	Gly	Ser	Arg	Ala	His 265	Ser	Ser	His	Leu	Lys 270	Ser	Lys	
			Lys	Gly	Gln 275	Ser	Thr	Ser	Arg	His 280	Lys	Lys	Leu	Met	Phe 285	Lys	Thr	Glu	
THE HEAT ALL ALL THE	35		Gly	Pro 290	Asp	Ser	Asp												
źa.	40	(2)	INFO	RMAT:	ION I	FOR	SEQ :	ID N	0:3:										
			(i)		) LEI	NGTH	ARAC' : 20 nucl	66 b	ase ]	pair	S								
	45			(C	) ST	RAND:	EDNE	ss:	sing:										
			(xi)	SEQ	UENC:	E DE	SCRI	PTIO	N: S	EQ I	D NO	:3:							
	50	CAAA	ACTT.	AC C	AGGG	CAAC	T AT	GGCT	TCCA	CCT	GGGC	TTC	CTGC.	AGTC	TG G	GACA	GCCA	A	60
	50	GTCI	GTTA	TG T	GCAC	GTAC'	T CT	CCTC	CCCT	CAA	TAAG	CTA	TTCT	GCCA	GC T	GGCG	AAGA	С	120

Arg His Ser Val Val Val Pro Tyr Glu Pro Pro Glu Val Gly Ser Asp

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15	(2) INFO	ITAMS	ON F	FOR S	SEQ ]	D NO	D:4:										
20	(i)	(B)	LEN TYP STF	E CHA NGTH: PE: & RANDE POLOG	293 minc EDNES	ami aci	ino a id singl	acids	3								
25	(xi)	SEQU	JENCI	E DES	CRII	PTIO	N: SI	EQ II	NO:	:4:							
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30	Gly	Thr	Ala	Lys 20	Ser	Val	Met	Cys	Thr 25	Tyr	Ser	Pro	Pro	Leu 30	Asn	Lys	
	Leu	Phe	Cys 35	Gln	Leu	Ala	Lys	Thr 40	Cys	Pro	Val	Gln	Leu 45	Trp	Val	Ser	
35	Ala	Thr 50	Pro	Pro	Ala	Gly	Ser 55	Arg	Val	Arg	Ala	Met 60	Ala	Ile	His	Lys	
40	Lys 65	Ser	Gln	His	Met	Thr 70	Gly	Val	Val	Arg	Arg 75	Cys	Pro	His	His	Glu 80	
40	Arg	Cys	Ser	Asp	Gly 85	Asp	Gly	Leu	Ala	Pro 90	Pro	Gln	His	Leu	Ile 95	Arg	
45	Val	Glu	Gly	Asn 100	Leu	Tyr	Pro	Glu	Tyr 105	Leu	Glu	Asp	Arg	Gln 110	Thr	Phe	
	Arg	His	Ser 115	Val	Val	Val	Pro	Tyr 120	Glu	Pro	Pro	Glu	Ala 125	Gly	Ser	Glu	

Tyr Thr Thr Ile His Tyr Lys Tyr Ile Cys Asn Ser Ser Cys Met Gly

	10		Glu	Val	Leu 195	Cys	Pro	Glu	Leu	Pro 200	Pro	Gly	Ser	Ala	Lys 205	Arg	Ala	Leu
	15		Pro	Thr 210	Cys	Thr	Ser	Ala	Ser 215	Pro	Pro	Gln	Lys	Lys 220	Lys	Pro	Leu	Asp
	13		Gly 225	Glu	Tyr	Phe	Thr	Leu 230	Lys	Ile	Arg	Gly	Arg 235	Leu	Arg	Phe	Glu	Met 240
	20		Phe	Arg	Glu	Leu	Asn 245	Glu	Ala	Leu	Glu	Leu 250	Lys	Asp	Ala	His	Ala 255	Thr
The House			Glu	Glu	Ser	Gly 260	Asp	Ser	Arg	Ala	His 265	Ser	Ser	Tyr	Leu	Lys 270	Ser	Lys
Hall and W.	25		Lys	Gly	Gln 275	Ser	Thr	Ser	Arg	His 280	Lys	Lys	Thr	Met	Val 285	Lys	Lys	Val
The first with their first factor and with	30		Gly	Pro 290	Asp	Ser	Asp											
An Hu		(2)	INFO	RMAT:	ION I	FOR S	SEQ	ID N	0:5:									
	35	(2) INFORMATION FOR SEQ ID NO:5:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 17 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear																
	40		(xi)	SEQ	UENC	E DE:	SCRI	PTIO	N: S	EQ I	D NO	:5:						
		ACTGCCCAAC AACACCA																

(2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single (D) TOPOLOGY: linear

Gly Met Asn Arg Arg Pro Ile Leu Thr Ile Ile Thr Leu Glu Asp Ser

Ser Gly Asn Leu Leu Gly Arg Asn Ser Phe Glu Val Arg Val Cys Ala

Cys Pro Gly Arg Asp Arg Thr Glu Glu Asn Phe Arg Lys Lys 

	GCCACGCCCA CACATTT	17
5	(2) INFORMATION FOR SEQ ID NO:7:	
10	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
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20	(2) INFORMATION FOR SEQ ID NO:8:  (i) SEQUENCE CHARACTERISTICS:	
25	(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
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35	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 18 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
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The state of the s

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

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5	(2) INFORMATION FOR SEQ ID NO:11:	
10	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
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	(2) INFORMATION FOR SEQ ID NO:13:	
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40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

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## What is Claimed is:

- 1. A method for treating a subject with a malignancy comprising the steps of:
- 5 (a) providing an expression construct comprising a promoter functional in eukaryotic cells and a polynucleotide encoding a *p53*, wherein said polynucleotide is positioned sense to and under the control of said promoter; and
  - (b) contacting said expression construct with a tumor cell in vivo.
- 10 2. The method of claim 1, wherein said malignancy is a squamous cell carcinoma.
  - 3. The method of claim 1, wherein the endogenous p53 of said tumor cell is mutated.
- 15 4. The method of claim 1, wherein the endogenous p53 of said tumor cell is wild-type.
  - 5. The method of claim 1, wherein said expression construct is a viral vector.
  - 6. The method of claim 5, wherein said viral vector is selected from the group consisting of a retroviral vector, an adenoviral vector and an adeno-associated viral vector.
  - 7. The method of claim 6, wherein said viral vector is a replication-deficient adenoviral vector.
  - 8. The method of claim 7, wherein said replication-deficient adenoviral vector is lacking at least a portion of the E1-region.
    - 9. The method of claim 8, wherein said promoter is a CMV IE promoter.
    - 10. The method of claim 1, wherein said subject is a human.
    - 11. The method of claim 7, wherein step (b) is repeated at least once.
- The method of claim 11, wherein said tumor is resected following a repeated contacting, and an additional contacting is effected subsequent to the resection.
  - 13. The method of claim 12, wherein said expression vector is contacted in a volume of about 3 ml. to about 10 ml.
  - 14. The method of claim 11, wherein the amount of adenovirus administered in each contacting is between about  $10^7$  and  $10^{12}$  pfu.

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- 15. The method of claim 1, wherein said contacting is via intratumoral injection.
- 16. The method of claim 1, wherein said contacting is via injection into a natural or artificial body cavity.
  - 17. The method of claim 16, wherein said injection comprises continuous perfusion of said natural or artificial body cavity.
- 10 18. The method of claim 16, wherein said contacting is via injection into an artificial body cavity resulting from tumor excision.
  - 19. The method of claim 1, wherein the p53-encoding polynucleotide is tagged so that expression of p53 from said expression vector can be detected.
    - 20. The method of claim 19, wherein the tag is a continuous epitope.
  - 21. A method for determining the effectiveness of a therapy on microscopic residual cancer comprising:
    - (a) providing a rodent with an incision into subcutaneous tissue;
    - (b) seeding said incision with tumor cells;
    - (c) treating said rodent with a therapeutic regimen; and
    - (d) assessing the impact of said regimen on the development of tumors.
  - 22. The method of claim 21, wherein said incision is sealed following step (b) and prior to step (c).
  - 23. The method of claim 22, wherein said therapeutic regimen comprises introduction of a therapeutic composition into said incision, said incision being reopened after sealing and resealed after introduction of said therapeutic composition.
  - 24. The method of claim 23, wherein said therapeutic composition comprises an expression construct comprising a promoter functional in eukaryotic cells and a polynucleotide encoding a p53, wherein said polynucleotide is positioned sense to and under the control of said promoter.
  - 25. The method of claim 24, wherein said expression construct is a replication-deficient adenovirus and said promoter is a CMV IE promoter.

## **ABSTRACT**

Methods for the treatment of squamous cell carcinoma using a p53-expressing viral vector are disclosed. In particular embodiments, the vector is a replication-deficient adenovirus.

In addition, there are provided methods for examining the development and treatment of microscopic residual disease in the context of post-surgical environments and in body cavities.

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#### SEQUENCE LISTING

- (1) GENERAL INFORMATION:
  - (i) APPLICANT: Clayman, Gary L.
  - (ii) TITLE OF INVENTION: Methods and Compositions for the Diagnosis and Treatment of Cancer
  - (iii) NUMBER OF SEQUENCES: 14
  - (iv) CORRESPONDENCE ADDRESS:
    - (A) ADDRESSEE: Arnold, White and Durkee
    - (B) STREET: P.O. Box 4433
    - (C) CITY: Houston
    - (D) STATE: TX
    - (E) COUNTRY: USA
    - (F) ZIP: 77210-4433
    - (v) COMPUTER READABLE FORM:
      - (A) MEDIUM TYPE: Floppy disk
      - (B) COMPUTER: IBM PC compatible
      - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
      - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
  - (vi) CURRENT APPLICATION DATA:
    - (A) APPLICATION NUMBER: UNKNOWN
    - (B) FILING DATE: CONCURRENTLY HEREWITH
    - (C) CLASSIFICATION: UNKNOWN
  - (viii) ATTORNEY/AGENT INFORMATION:
    - (A) NAME: Highlander, Steven L.
    - (B) REGISTRATION NUMBER: 37,642
    - (C) REFERENCE/DOCKET NUMBER: INGN:022
    - (ix) TELECOMMUNICATION INFORMATION:
      - (A) TELEPHONE: (512) 418-3000
      - (B) TELEFAX: (512) 474-7577
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 2066 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

CAAAACCTAC CAGGGCAGCT ACGGTTTCCG TCTGGGCTTC TTGCATTCTG GGACAGCCAA 60
GTCTGTGACT TGCACGTACT CCCCTGCCCT CAACAAGATG TTTTGCCAAC TGGCCAAGAC 120

CTGCCCTGTG CAGCTGTGGG TTGATTCCAC ACCCCCGCCC GGCACCCGCG TCCGCGCCAT 180 GGCCATCTAC AAGCAGTCAC AGCACATGAC GGAGGTTGTG AGGCGCTGCC CCCACCATGA 240 GCGCTGCTCA GATAGCGATG GTCTGGCCCC TCCTCAGCAT CTTATCCGAG TGGAAGGAAA 300 TTTGCGTGTG GAGTATTTGG ATGACAGAAA CACTTTTCGA CATAGTGTGG TGGTGCCCTA 360 TGAGCCGCCT GAGGTTGGCT CTGACTGTAC CACCATCCAC TACAACTACA TGTGTAACAG 420 TTCCTGCATG GGCGGCATGA ACCGGAGGCC CATCCTCACC ATCATCACAC TGGAAGACTC 480 CAGTGGTAAT CTACTGGGAC GGAACAGCTT TGAGGTGCGT GTTTGTGCCT GTCCTGGGAG 540 AGACCGGCGC ACAGAGGAAG AGAATCTCCG CAAGAAAGGG GAGCCTCACC ACGAGCTGCC 600 660 CCCAGGGAGC ACTAAGCGAG CACTGCCCAA CAACACCAGC TCCTCTCCCC AGCCAAAGAA GAAACCACTG GATGGAGAAT ATTTCACCCT TCAGATCCGT GGGCGTGAGC GCTTCGAGAT 720 GTTCCGAGAG CTGAATGAGG CCTTGGAACT CAAGGATGCC CAGGCTGGGA AGGAGCCAGG 780 GGGGAGCAGG GCTCACTCCA GCCACCTGAA GTCCAAAAAG GGTCAGTCTA CCTCCCGCCA 840 900 TAAAAAACTC ATGTTCAAGA CAGAAGGGCC TGACTCAGAC TGACATTCTC CACTTCTTGT TCCCCACTGA CAGCCTCCCA CCCCCATCTC TCCCTCCCCT GCGATTTTGG GTTTTGGGTC 960 TTTGAACCCT TGCTTGCAAT AGGTGTGCGT CAGAAGCACC CAGGACTTCC ATTTGCTTTG 1020 TCCCGGGGCT CCACTGAACA AGTTGGCCTG CACTGGTGTT TTGTTGTGGG GAGGAGGATG 1080 GGGAGTAGGA CATACCAGCT TAGATTTTAA GGTTTTTACT GTGAGGGATG TTTGGGAGAT 1140 GTAAGAATG TTCTTGCAGT TAAGGGTTAG TTTACAATCA GCCACATTCT AGGTAGGGGC 1200 CCACTTCACC GTACTAACCA GGGAAGCTGT CCCTCACTGT TGAATTTTCT CTAACTTCAA 1260 GGCCCATATC TGTGAAATGC TGGCATTTGC ACCTACCTCA CAGAGTGCAT TGTGAGGGTT 1320 AATGAAATAA TGTACATCTG GCCTTGAAAC CACCTTTTAT TACATGGGGT CTAGAACTTG 1380 ACCCCCTTGA GGGTGCTTGT TCCCTCTCCC TGTTGGTCGG TGGGTTGGTA GTTTCTACAG 1440 TTGGGCAGCT GGTTAGGTAG AGGGAGTTGT CAAGTCTCTG CTGGCCCAGC CAAACCCTGT 1500 CTGACAACCT CTTGGTGAAC CTTAGATCCT AAAAGGAAAT GTCACCCCAT CCCACACCCT 1560 GGAGGATTTC ATCTCTTGTA TAGATGATCT GGATCCACCA AGACTTGTTT TAGCTCAGGG 1620 TCCAATTTCT TTTTTCTTTT TTTTTTTTT TTTCTTTTC TTTGAGACTG GGTCTCTTTG 1680 TTGCCCCAGG CTGGAGTGGA GTGGCGTGAT CTGGCTTACT GCAGCCTTTG CCTCCCCGGC 1740 TCGAGCAGTC CTGCCTCAGC CTCCGGAGTA GCTGGGACCA CAGGTTCATG CCACCATGGC 1800

CAGCCAACTT TTGCATGTTT TGTAGAGATG GGGTCTCACA GTGTTGCCCA GGCTGGTCTC 1860

AAACTCCTGG GCTCAGGCGA TCCACCTGTC TCAGCCTCCC AGAGTGCTGG GATTACAATT 1920

GTGAGCCACC ACGTCCAGCT GGAAGGGTCA ACATCTTTA CATTCTGCAA GCACATCTGC 1980

ATTTTCACCC CACCCTTCCC CTCTTCTCCC TTTTTATATC CCATTTTAT ATCGATCTCT 2040

TATTTTACAA TAAAACTTTG CTGCCA 2066

#### (2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 293 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Lys Thr Tyr Gln Gly Ser Tyr Gly Phe Arg Leu Gly Phe Leu His Ser 1 5 10 15

Gly Thr Ala Lys Ser Val Thr Cys Thr Tyr Ser Pro Ala Leu Asn Lys 20 25 30

Met Phe Cys Gln Leu Ala Lys Thr Cys Pro Val Gln Leu Trp Val Asp 35 40 45

Ser Thr Pro Pro Pro Gly Thr Arg Val Arg Ala Met Ala Ile Tyr Lys 50 55 60

Gln Ser Gln His Met Thr Glu Val Val Arg Arg Cys Pro His His Glu 65 70 75 80

Arg Cys Ser Asp Ser Asp Gly Leu Ala Pro Pro Gln His Leu Ile Arg 85 90 95

Val Glu Gly Asn Leu Arg Val Glu Tyr Leu Asp Asp Arg Asn Thr Phe 100 105 110

Arg His Ser Val Val Val Pro Tyr Glu Pro Pro Glu Val Gly Ser Asp 115 120 125

Cys Thr Thr Ile His Tyr Asn Tyr Met Cys Asn Ser Ser Cys Met Gly 130 135 140

Gly Met Asn Arg Arg Pro Ile Leu Thr Ile Ile Thr Leu Glu Asp Ser 145 150 155 160

Ser Gly Asn Leu Leu Gly Arg Asn Ser Phe Glu Val Arg Val Cys Ala

165 170 175

Cys Pro Gly Arg Asp Arg Thr Glu Glu Glu Asn Leu Arg Lys Lys 180 185 190

Gly Glu Pro His His Glu Leu Pro Pro Gly Ser Thr Lys Arg Ala Leu 195 200 205

Pro Asn Asn Thr Ser Ser Ser Pro Gln Pro Lys Lys Pro Leu Asp 210 215 220

Gly Glu Tyr Phe Thr Leu Gln Ile Arg Gly Arg Glu Arg Phe Glu Met 225 230 235 240

Phe Arg Glu Leu Asn Glu Ala Leu Glu Leu Lys Asp Ala Gln Ala Gly 245 250 255

Lys Glu Pro Gly Gly Ser Arg Ala His Ser Ser His Leu Lys Ser Lys 260 265 270

Lys Gly Gln Ser Thr Ser Arg His Lys Lys Leu Met Phe Lys Thr Glu 275 280 285

Gly Pro Asp Ser Asp 290

#### (2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2066 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

CAAAACTTAC	CAGGGCAACT	ATGGCTTCCA	CCTGGGCTTC	CTGCAGTCTG	GGACAGCCAA	60
GTCTGTTATG	TGCACGTACT	CTCCTCCCCT	CAATAAGCTA	TTCTGCCAGC	TGGCGAAGAC	120
GTGCCCTGTG	CAGTTGTGGG	TCAGCGCCAC	ACCTCCAGCT	GGGAGCCGTG	TCCGCGCCAT	180
GGCCATCCAC	AAGAAGTCAC	AGCACTTGAC	GGGGGTCGTG	AGACGCTGCC	CCCACCATGA	240
GCGCTGCTCC	GATGGTGATG	GCCTGGCTCC	TCCCCAGCAT	CTTATCCGGG	TGGAAGGAAA	300
TTTGTATCCC	GAGTATCTGG	AAGACAGGCA	GACTTTTCGC	CACAGCGTGG	TGGTACCTTA	360
TGAGCCACCC	GAGGCCGGCT	CTGAGTATAC	CACCATCCAC	TACAAGTACA	TTTGTAATAG	420
CTCCTGCATG	GGGGGCATGA	ACCGCCGACC	TATCCTTACC	ATCATCACAC	TGGAAGACTC	480
CAGTGGGAAC	CTTCTGGGAC	GGGACAGCTT	TGAGGTTCGT	GTTTGTGCCT	GCCCTGGGAG	540

AGACCGCCGT ACAGAAGAAG AAAATTTCCG CAAAAAGGAA GTCCTTTGCC CTGAACTGCC 600 CCCAGGGAGC GCAAAGAGA CGCTGCCCAC CTGCACAAGC GCCTCTCCCC CGCAAAAGAA 660 AAAACCACTT GATGGAGAGT ATTTCACCCT CAAGATCCGC GGGCGTAAAC GCTTCGAGAT 720 GTTCCGGGAG CTGAATGAGG CCTTAGAGTT AAAGGATGCC CATGCTACAG AGGAGTCTGG 780 AGACAGCAGG GCTCACTCCA GCTACCTGAA GACCAAGAAG GGCCAGTCTA CTTCCCGCCA 840 TAAAAAAACA ATGGTCAAGA AAGTGGGGCC TGACTCAGAC TGACATTCTC CACTTCTTGT 900 TCCCCACTGA CAGCCTCCCA CCCCCATCTC TCCCTCCCCT GCCTTTTGGG TTTTGGGTCT 960 TTGAACCCTT GCTTGCAATA GGTGTGCGTC AGAAGCACCC AGGACTTCCA TTTGCTTTGT 1020 CCCGGGGCTC CACTGAACAA GTTGGCCTGC ACTGGTGTTT TGTTGTGGGG AGGAGGATGG 1080 GGAGTAGGAC ATACCAGCTT AGATTTTAAG GTTTTTACTG TGAGGGATGT TTGGGAGATG 1140 TAAGAAATGT TCTTGCAGTT AAGGGTTAGT TTACAATCAG CCACATTCTA GGTAGGGGCC 1200 CACTTCACCG TACTAACCAG GGAAGCTGTC CCTCACTGTT GAATTTTCTC TAACTTCAAG 1260 GCCCATATCT GTGAAATGCT GGCATTTGCA CCTACCTCAC AGAGTGCATT GTGAGGGTTA 1320 ATGAAATAAT GTACATCTGG CCTTGAAACC ACCTTTTATT ACATGGGGTC TAGATGACCC 1380 CCTTGAGGTG CTTGTTCCCT CTCCCTGTTG GTCGGTGGGT TGGTAGTTTC TACAGTTGGG 1440 CAGCTGGTTA GGTTGAGGTA GTTGTCAGGT CTCTGCTGGC CCAGCGAAAT TCTATCCAGC 1500 CAGTTGTTGG ACCCTGGCAC CTCAAATGAA ATCTCACCCT ACCCCACACC CTGTAAGATT 1560 1620 CTATCTCTTG TATAGATGAT CTGGATCCAC CAAGACTTGT TTTAGCTCAG GGTCCAATTT CTTTTTCTT TTTTTTTTT TTTTTCTTTT TCTTTGAGAC TGGGTCTCTT TGTTGCCCCA 1680 GGCTGGAGTG GAGTGGCGTG ATCTGGCTTA CTGCAGCCTT TGCCTCCCCG GCTCGAGCAG 1740 TCCTGCCTCA GCCTCCGGAG TAGCTGGGAC CACAGGTTCA TGCCACCATG GCCAGCCAAC 1800 TTTTGCATGT TTTGTAGAGA TGGGGTCTCA CAGTGTTGCC CAGGCTGGTC TCAAACTCCT 1860 GGGCTCAGGC GATCCACCTG TCTCAGCCTC CCAGAGTGCT GGGATTACAA TTGTGAGCCA 1920 CCACGTCCAG CTGGAAGGGC CTACTTTCCT TCCATTCTGC AAAGCCCTGC TGCATTTATC 1980 CACCCCACCC TCCACCTGTC TCCCTCTTTT TTTCTTACCC CTTTTTATAT ATCAATTTCT 2040 2066 TATTTTACAA TAAAATTTTG TTATCA

#### (2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 293 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Lys Thr Tyr Gln Gly Asn Tyr Gly Phe His Leu Gly Phe Leu Gln Ser 1 5 10 15

Gly Thr Ala Lys Ser Val Met Cys Thr Tyr Ser Pro Pro Leu Asn Lys 20 25 30

Leu Phe Cys Gln Leu Ala Lys Thr Cys Pro Val Gln Leu Trp Val Ser 35 40 45

Ala Thr Pro Pro Ala Gly Ser Arg Val Arg Ala Met Ala Ile His Lys
50 55 60

Lys Ser Gln His Met Thr Gly Val Val Arg Arg Cys Pro His His Glu 65 70 75 80

Arg Cys Ser Asp Gly Asp Gly Leu Ala Pro Pro Gln His Leu Ile Arg 85 90 95

Val Glu Gly Asn Leu Tyr Pro Glu Tyr Leu Glu Asp Arg Gln Thr Phe
100 105 110

Arg His Ser Val Val Val Pro Tyr Glu Pro Pro Glu Ala Gly Ser Glu
115 120 125

Tyr Thr Thr Ile His Tyr Lys Tyr Ile Cys Asn Ser Ser Cys Met Gly 130 135 140

Gly Met Asn Arg Arg Pro Ile Leu Thr Ile Ile Thr Leu Glu Asp Ser 145 150 155 160

Ser Gly Asn Leu Leu Gly Arg Asn Ser Phe Glu Val Arg Val Cys Ala 165 170 175

Cys Pro Gly Arg Asp Arg Arg Thr Glu Glu Glu Asn Phe Arg Lys Lys
180 185 190

Glu Val Leu Cys Pro Glu Leu Pro Pro Gly Ser Ala Lys Arg Ala Leu 195 200 205

Pro Thr Cys Thr Ser Ala Ser Pro Pro Gln Lys Lys Pro Leu Asp 210 215 220

Gly Glu Tyr Phe Thr Leu Lys Ile Arg Gly Arg Leu Arg Phe Glu Met 225 230 235 240

	Phe	Arg	Glu	Leu	Asn 245	Glu	Ala	Leu	Glu	Leu 250	Lys	Asp	Ala	His	Ala 255	Thr	
	Glu	Glu	Ser	Gly 260	Asp	Ser	Arg	Ala	His 265	Ser	Ser	Tyr	Leu	Lys 270	Ser	Lys	
	Lys	Gly	Gln 275	Ser	Thr	Ser	Arg	His 280	Lys	Lys	Thr	Met	Val 285	Lys	Lys	Val	
	Gly	Pro 290	Asp	Ser	Asp												
(2)	INFO	RMAT:	ION I	FOR a	SEQ :	ID N	0:5:										
	(i)	(B)	) LE ) TY: ) ST:	NGTH PE: : RAND:	ARAC' : 17 nucl EDNE GY:	base eic a	e pa acid sing	irs									
	(xi)	SEQ	UENC:	E DE	SCRI	PTIO	N: S	EQ I	D NO	:5:							
ACTG	CCCA	AC A	ACAC	CA													17
(2)	INFO	RMAT	ION :	FOR	SEQ	ID N	0:6:										
	(i)	(B (C	) LE ) TY ) ST	NGTH PE: RAND	ARAC : 17 nucl EDNE GY:	bas eic SS:	e pa acid sing	irs									
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	:6:							
GCC	ACGCC	CA C	ACAT	TT													17
(2)	INFO	RMAT	ION	FOR	SEQ	ID N	0:7:										
	(i)	(B	) LE ) TY ) ST	NGTH PE: RAND	ARAC : 20 nucl EDNE	bas eic SS:	e pa acid sing	irs l									
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NC	):7:							
GCC.	rgrcc	TG G	GAGA	.GACC	:G												20

(2) INFORMATION FOR SEQ ID NO:8:

(1)	(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:8:	
CCCTTAAG	CC ACGCCCACAC	20
(2) INFO	RMATION FOR SEQ ID NO:9:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:9:	
CACTGCCC	AA CAACACCA	18
(2) INFO	RMATION FOR SEQ ID NO:10:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 17 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:10:	
GCCACGCC	CA CACATTT	17
(2) INFC	ORMATION FOR SEQ ID NO:11:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:11:	
GGTGCATI	TGG AACGCGGATT	20
(2) INFO	ORMATION FOR SEQ ID NO:12:	

(i) SEQUENCE CHARACTERISTICS:

Strang.	HH
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7	#
walke than	H Hr.
The court of the court	K., H., Hr.
The state of the state	A.A. M M Pr
walke than	A.A. M M Pr

(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
GGGGACAGAA CGTTGTTTC	2
(2) INFORMATION FOR SEQ ID NO:13:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
ACGGATTTGG TCGTATTGGG	2
(2) INFORMATION FOR SEQ ID NO:14:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
TGATTTTGGA GGGATCTCGC	2

(A) LENGTH: 20 base pairs(B) TYPE: nucleic acid

FIG 1

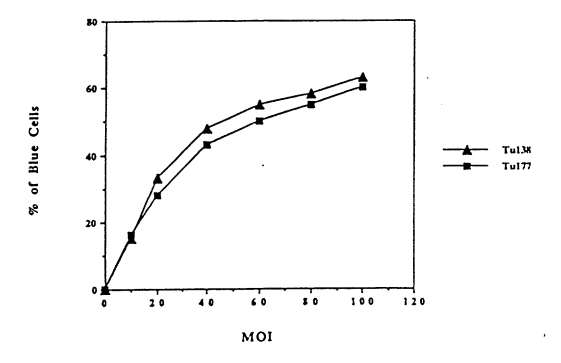


FIG 2A

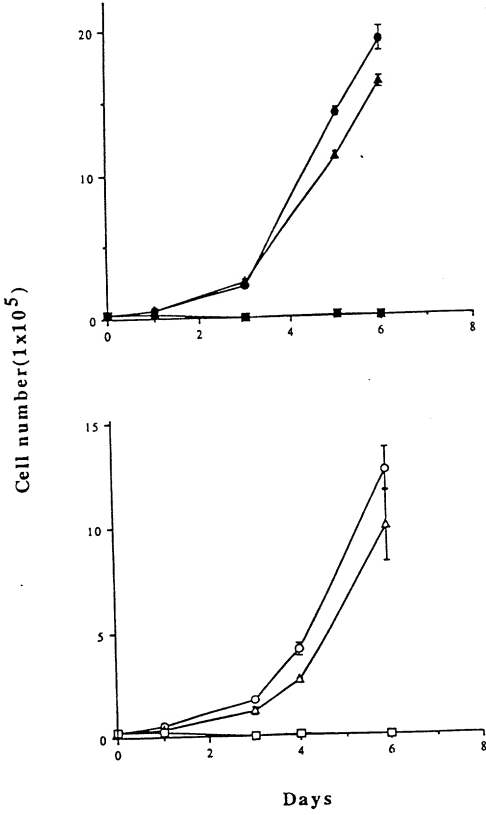


FIG 2B

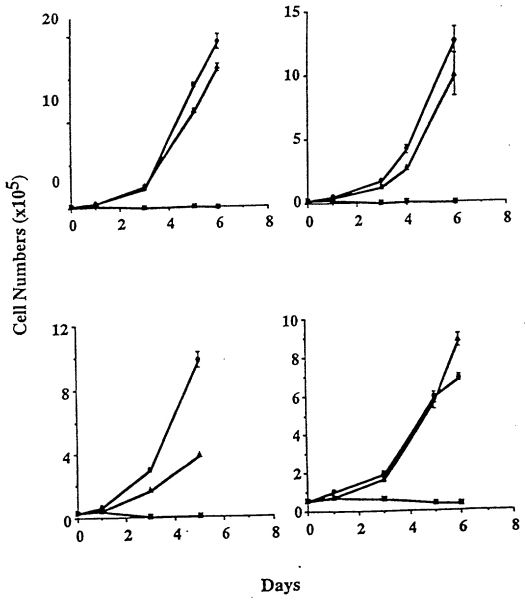
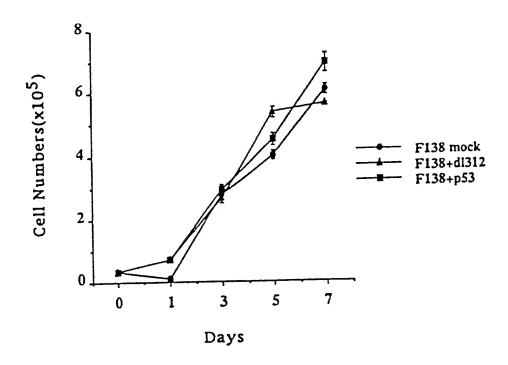


FIG 3C

FIG 3D

FIG 4



# FIG 5A

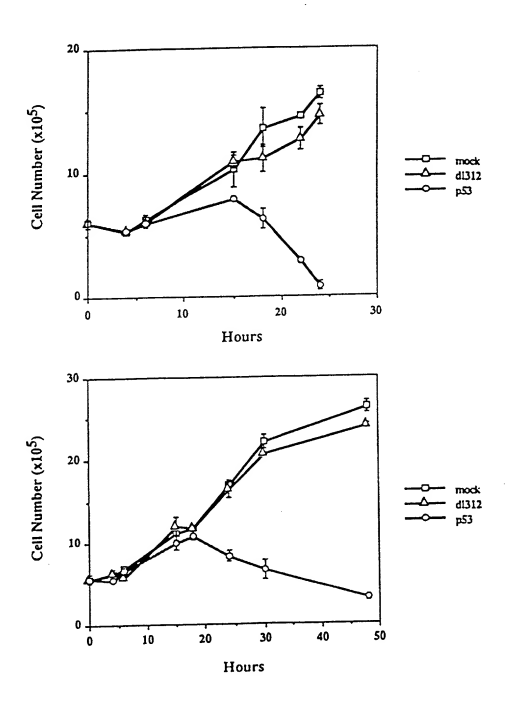


FIG 5B

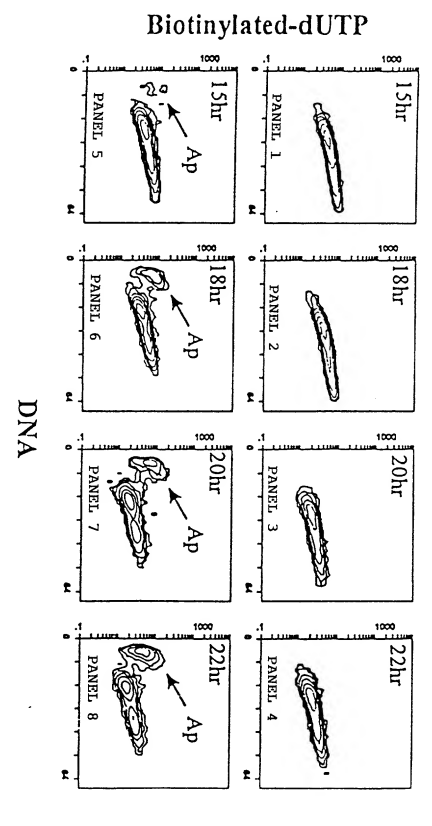


FIG 6A

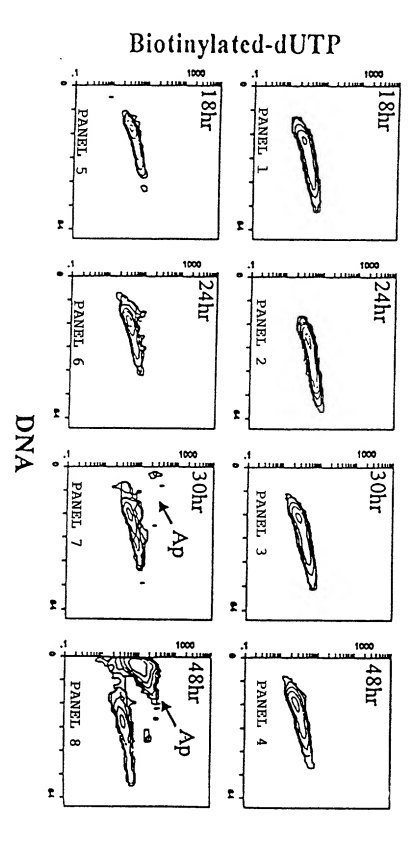


FIG 6B

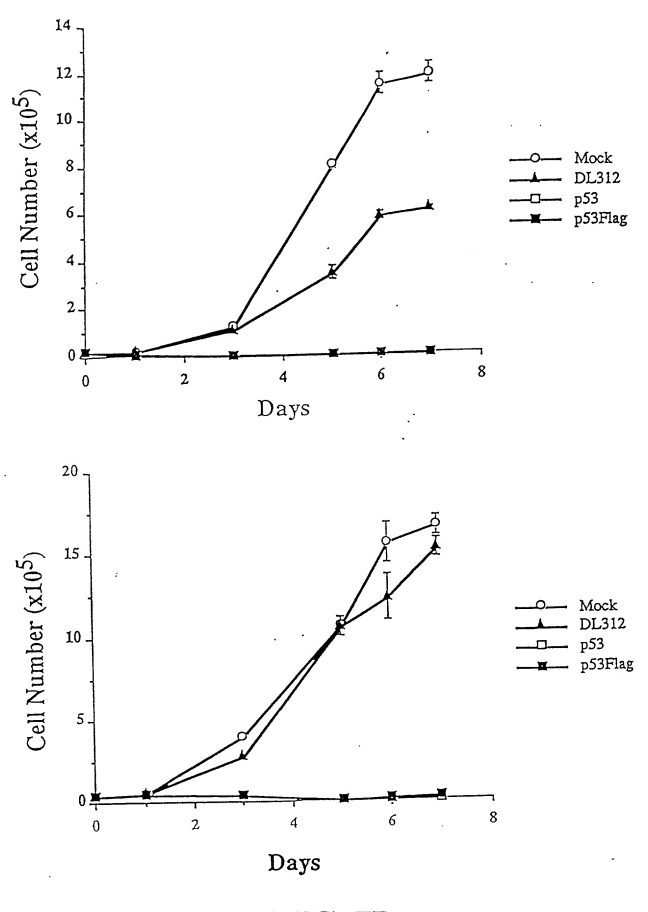


FIG 7B





#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of: Gary L. Clayman § § Examiner: Unknown Serial No.: 08/758,033 Group Art Unit: Unknown Filed: November 27, 1996 Atty. Dkt: INGN:041/HYL For: METHODS AND COMPOSITIONS FOR THE DIAGNOSIS AND TREATMENT OF CANCER

#### DECLARATION CLAIMING SMALL ENTITY STATUS 37 C.F.R. §§ 1.9(f) and 1.27(d) - NONPROFIT ORGANIZATION

Assistant Commissioner for Patents Washington, D.C. 20231

Sir:

I hereby declare that I am an official empowered to act on behalf of the nonprofit organization identified below:

Name of Organization:

Board of Regents,

The University of Texas System

Address of Organization: 201 West 7th Street

Austin, Texas 78701

The type of organization is a university.

I hereby declare that the organization identified above qualifies as a nonprofit organization as defined in 37 C.F.R. § 1.9(e) (1), and thus is a "small entity" as defined in § 1.9(f), for purposes of paying reduced fees under Sections 41(a) and (b) of Title 35, United States Code, with regard to the abovereferenced application.

I hereby declare that exclusive rights to the invention have been conveyed to and remain with the organization, with respect to the above-referenced invention, nor have I assigned, granted, conveyed or licensed and am under no obligation under contract or law to assign, grant, convey or license, any rights in the invention to any person who could not be classified as an independent inventor under 37 CFR § 1.9(c) if that person had made the invention, or to any concern which would not qualify as a small business concern under 37 CFR § 1.9(d) or a nonprofit organization under 37 CFR § 1.9(e), with the exception that the Government may have rights in the invention pursuant to a funding agreement under 35 U.S.C. § 202(c)(4): None

I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate.

There of the fleet faith from the fleet fleet bank Marie He He M I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.

BOARD OF RECENTS,

THE UNIVERSITY OF TEXAS SYSTEM

By:

Name: Ray Farabee Title: Vice Chancellor and General Counsel

Date: 3/10/97

#### DECLARATION

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or the below named inventors are the original, first and joint inventors (if plural names are listed below) of the subject matter which s claimed and for which a patent is sought on the invention entitled METHODS AND COMPOSITIONS FOR THE DIAGNOSIS AND TREATMENT OF CANCER, the Specification of which:

is attached hereto.  $\overline{X}$  was filed on November 27, 1996 as Application Serial No. 08/758,033.

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims.

I acknowledge the duty to disclose to the Patent and Trademark Office all information known to me to be material to patentability of the subject matter claimed in this application, as "materiality" is defined in Title 37, Code of Federal Regulations, § 1.56.

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose all information known to me to be material to patentability of the subject matter claimed in this application, as "materiality" is defined in Title 37, Code of Federal Regulations, § 1.56, which become available between the filing date of the prior application and the national or PCT international filing date of this application:

60/007,810 November 30, 1995 Pending (Application Serial No.) (Filing Date) (Status)

I hereby direct that all correspondence and telephone calls be addressed to Steven L. Highlander, Arnold, White & Durkee, P.O. Box 4433, Houston, Texas 77210 (512) 418-3000.

I hereby declare that all statements made of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Inventor's Full Name: Gary L. Clayman

(First) (Initial) (Last)

Inventor's Signature:

Date: Fw2+4,1997 Country of Citizenship: USA

Resident Address: MSS | MSS

(if different from residence address)

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

ELECTION UNDER 37 C.F.R. §§ 3.71
AND 3.73 AND POWER OF ATTORNEY

Assistant Commissioner for Patents Washington, D.C. 20231

Sir:

The undersigned, being Assignee of record of the entire interest in the above-identified application by virtue of an assignment recorded in the United States Patent and Trademark Office as set forth below, hereby elects, under 37 C.F.R. § 3.71, to prosecute the application to the exclusion of the inventor.

The Assignee hereby revokes any previous Powers of Attorney and appoints Louis T. Pirkey, Reg. No. 22,393; J. Paul Williamson, Reg. No. 29,600; Daniel S. Hodgins, Reg. No. 31,026; David L. Parker, Reg. No. 32,165; Barbara S. Kitchell, Reg. No. 33,928; Mark B. Wilson, Reg. No. 37,259; Gary J. Sertich, Reg. No. 34,430; Steven L. Highlander, Reg. No. 37,642; Timothy S. Corder, Reg. No. 38,414; Adam V. Floyd, Reg. No. 39,192; and Shelley P.M. Fussey, Reg. No. 39,458; each an attorney or agent of the firm of ARNOLD, WHITE & DURKEE, as its attorney or agent for so long as they remain with such firm, with full power of substitution and revocation, to prosecute the application, to make alterations and amendments therein, to transact all business in the Patent and Trademark Office in connection therewith, and to receive any Letters Patent, and for one year after issuance of such Letters Patent to file any request for a certificate of correction that may be deemed appropriate.

Pursuant to 37 C.F.R. § 3.73, the undersigned has reviewed the evidentiary documents, specifically the Assignment to Board of Regents, The University of Texas System, referenced below, and certifies that to the best of my knowledge and belief, title remains in the name of the Assignee.

Please direct all communications as follows:

Steven L. Highlander ARNOLD, WHITE & DURKEE P.O. Box 4433 Houston, Texas 77210-4433 (512) 418-3000

ASSIGNEE:

BOARD OF RECENTS, THE

UNIVERSITY OF TEXAS SYSTEM

Name: Ray Farabee

Title: Vice Chancellor and General Counsel

Date: 3/11/97
ASSIGNMENT: Concurrently filed